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UTILITY

PATENT APPLICATION

Approved for use through 10/31/2002 OMB 0651-0032

Patent and Trademark Office US DEPARTMENT OF COMMERCE are required to respond to a collection of information unless it displays a valid OMB control number. M8540/248465 Attorney Docket No. First Named Inventor Gary Anthony Jubb

TRA	NSMITTAL	or Applicati	on Identifier Gary Antirony Subb			
	rovisional applications under CFR 1.53(b))	Title	Saline Soluble Ir		le Inorganic Fibres	
		Ехр	oress Mail Label No. EL568948202US			
APPL	APPLICATION ELEMENTS			TO: Assistai	nt Director for Pa	atents
	cerning utility patent application contents			Box Pat	ent Application gton, D.C 2023	· 1
1. Fee Transing (Submit an original Submit and origi	nittal Form (e.g. PTO/SB/17 ginal, and a duplicate for fee processing) laims small entity status. R 1.27 In Total Pages gement as set forth below) ive title of the Invention eferences to Related Application eferences to Related Application ent Regarding Fed sponsore ce to sequence listing, a table er program listing appendix und of the Invention mmary of the Invention scription of the Drawings (if Description cof the Disclosure (35 U.S.C. 113) [Total Sheet claration [Total Pages ewly executed (original or compy from a prior application 63(d)) (for continuation/division 2. 17 completed) DELETION OF INVENTOR Signed statement attached deletin named in the prior application, See 1.63(d)(2) and 1.33(b). Data Sheet. See 37 CFR	ations d R & D ole, or a filed) ts] 05 02 oppy) (37 CFR sional with 8(S) g inventor(s) a 37 C.F.R.	Comp 8. Nucleotide (if applicable) a.	Washin COM or CD-R in dup puter Program (App e and/or Amino Acid, all necessary) emputer Readable Copecification Sequence CD-ROM or CD-F Paper statement verifying to CD-ROM or CD-F Paper of COMPANYING APP (COMPANYING APP) (COMP	property of a company of the company	e table or e Submission on:); or above copies PARTS document(s)) ower of Attorney opplicable) pies of IDS itations 03) t(s) awings er of Attorney
17. If a CONTINUING APPLICATION, check appropriate box and supply the requisite information below and in a preliminary amendment, or in an Application Data Sheet under 37 CFR 1.76: Continuation Divisional Continuation-in-part (CIP) of prior application No.: 09/262,378 Prior application information Examiner: K. Group Group/Art Unit: 1755 For CONTINUATION or DIVISIONAL APPS only: The entire disclosure of the prior application, from which an oath or declaration is supplied under Box 5b, is considered a part of the disclosure of the accompanying continuation or divisional application and is hereby incorporated by reference. The incorporation can only be relied upon when a portion has been inadvertently omitted from the submitted application parts. 18. CORRESPONDENCE ADDRESS						
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Name	John S. Pratt KILPATRICK STOCKTON	IIID		·····		
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Address	Suite 2800					
City	Atlanta		State	GA Zip	Code	30309
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FEE TRANSMITTAL	Complete if Known					
for FY 2000	Application Number	Divisional of 09/262,378	0			
Patent fees are subject to annual revision	Filing Date	October 24, 2000	4			
Small Entity payments must be supported by a small entity	First Named Inventor	Gary Anthony Jubb	.0			
statement, otherwise, large entity fees must be paid.	Group / Art Unit	1755	- 55 <u>-</u>			
See 37 C.F.R. §§ 1.27 and 1.28	Examiner Name	Group, K.	9			
TOTAL AMOUNT OF PAYMENT (\$)710.00	Attorney Docket Number	M8540/248465	92			

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Applicant claims small entity status. See 37 CFR 1,27	113	1,840*	113	1,840*		publication of SIR after	
2. Payment Enclosed:	115	110	215	55	Extension f	for reply within first month	
Check Credit Card Money Order Other	116	380	216	190	Extension for month	for reply within second	
FEE CALCULATION	117	870	217	435	Extension for	for reply within third month	,
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108 690 208 345 Reissue filing fee	141	1,210	241	605	Petition to r	revive - unintentional	
114 150 214 75 Provisional filing fee	142	1,210	242	605		e fee (or reissue)	
SUBTOTAL (1) (\$)710.00	143	430	243	215	Design issue		
2. EXTRA CLAIM FEES Extra Fee from Fee Paid	144	580	244	290	Plant issue f	fee	
Claims below	122	130	120	130	Petitions to	the Commissioner	
Total Claims 09 -20**= 00 x = .00	123	50	123	50		lated to provisional	
Independent Claims 01 -3**= 00 x = .00	126	240	126	240	Submission	s of Information Disclosure	,
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Signature					Date	121/00	

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicants:

Gary Anthony Jubb and Jean-Louis Martin

Serial No.:

Divisional of 09/262,378

Filing Date:

October 24, 2000

For:

SALINE SOLUBLE INORGANIC FIBRES

Box Patent Application Director of Patents and Trademarks Washington, D.C. 20231

Date: October 24, 2000

PRELIMINARY AMENDMENT

Sir:

Preliminary to any examination on the merits, Applicants respectfully submit the following amendments and remarks:

In the Specification

Kindly add the following paragraph immediately preceding line 1:

--This application is a divisional of U.S. Serial No. 09/262,378 filed March 4, 1999, now allowed, which is a continuation of U.S. Serial No. 08/899,005 filed July 23, 1997, now U.S. Patent No. 5,994,247 issued on November 30, 1999, which is a continuation of U.S. Serial No. 08/535,587 filed September 28, 1995, now abandoned, which is a continuation of U.S. Serial No. 08/039,086, filed April 9, 1993, now abandoned, which claims priority to International Application No. PCT/GB93/00085, filed January 15, 1993, which claims priority to GB 92 00993.5, filed January 17, 1992 and GB 92 24612.3, filed November 24, 1992--.

In the Claims

Please cancel claims 1-13 without prejudice or disclaimer to the subject matter thereof.

Please add the following new claims.

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Filed: October 24, 2000

PRELIMINARY AMENDMENT

--14. A method of insulating an article against temperatures which may on occasion exceed 900°C comprising:

disposing on, in, near or around the article thermal insulation which is a refractory insulating material having a maximum service temperature greater than 900°C and comprising vitreous fibers having a composition comprising SiO₂, CaO, MgO, and optionally Al₂O₃, wherein:

- (a) SiO_2 is present in an amount (1) greater than 58% by weight SiO_2 , if the amount of MgO in the composition is in the range 0 through 10 percent by weight; or (2) greater than the sum of (58 + 0.5) (weight percent of MgO 10) percent by weight SiO_2 , if the amount of MgO in the composition is greater than 10 percent by weight;
 - (b) an amount up to 42 percent by weight CaO;
 - (c) an amount up to 31.33 percent by weight MgO., and
 - (d) 0 to less than 3.97 percent by weight Al₂O₃;

wherein the refractory insulation material has a maximum service temperature greater than 900°C; the refractory insulation material has a shrinkage of less than 3.5 percent when exposed to a temperature of 1000°C for 24 hours, and has a shrinkage of less than 3.5 percent when exposed to a temperature of 800°C for 24 hours; and

wherein the refractory insulation material is essentially free of alkali metal oxide and boron oxide fluxing components.--

--15. The method of claim 14, wherein the amount of SiO₂ is less than 70 percent by weight. --

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PRELIMINARY AMENDMENT

- --16. The method of claim 14, wherein the vitreous fibers crystallize as wollastonite, pseudowollastonite, or a mixture thereof after exposure to a temperature of 1000°C, and wherein said wollastonite, pseudowollastonite, or mixture thereof comprises:
 - (a) 60 through 67 percent by weight SiO₂;
 - (b) 26 through 35 percent by weight CaO;
 - (c) 4 through 6 percent by weight MgO; and
 - (d) 0 through 3.5 percent by weight Al_2O_3 . --
 - --17. The method of claim 14, wherein the vitreous fibers comprise:
 - (a) a maximum of 71.24 percent by weight SiO₂;
 - (b) 4.46 through 34.49 percent by weight CaO;
 - (c) 1.71 through 22.31 percent by weight MgO; and
 - (d) 0 through 2.57 percent by weight Al_2O_3 . --
 - --18. The method of claim 17, wherein the vitreous fibers further comprise:
 - (e) 0 through 0.65 percent by weight Na₂O;
 - (f) 0 through 0.13 percent by weight K_2O ;
 - (g) 0.08 through 0.4 percent by weight Fe₂O₃; and
 - (h) 0 through 1.23 percent by weight ZrO_2 . --
- --19. The method of claim 14, further comprising forming said vitreous fibers into a bulk, blanket, block, or vacuum-formed form.--
- --20. The method of claim 14, wherein said insulated article is occasionally exposed to a temperature above around 1000°C.--

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- --21. The method of claim 19 wherein said vitreous fibers are formed into a needled blanket.--
- --22. The method of claim 14, wherein the vitreous fibers crystallize as diopside after exposure to a temperature of 1000°C, and wherein said diopside comprises:
 - (a) 59 through 64 percent by weight SiO₂;
 - (b) 19 through 23 percent by weight CaO;
 - (c) 14 through 17 percent by weight MgO; and
 - (d) 0 through 3.5 percent by weight Al_2O_3 .--

REMARKS

New claims 14-22 are fully supported by the specification as originally filed, and correspond to claims 23-28, 30, and 31 in parent application Serial No. 09/262,378. No new matter has been added.

Applicants respectfully submit that the claims are novel and nonobvious over Olds et al. (U.S. Patent No. 5,332,699), Olds et al. (WO 87/05007), or Karppinen et al., and that the rejections made by the Examiner over these references in the parent application should not be repeated.

The claims recite a process for insulating an article against temperatures that may exceed 900 °C. None of the above references teach insulating against such a high temperature, which is

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"Saline Soluble Inorganic Fibers"

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PRELIMINARY AMENDMENT

quite different from insulating against another, lower temperature. As a result, the references do

not anticipate the claimed invention. Moreover, there is no teaching or suggestion to use the

materials described in the above references to insulate against such high temperatures. In

particular, there is no teaching or suggestion that the materials described in the references would

be able to withstand temperatures that exceed 900 °C without shrinking. As the Examiner is no

doubt aware, even if a fiber can withstand high temperatures, if it shrinks substantially when

exposed to those temperatures, then its value as insulation is very low, since the shrinkage will

cause gaps in the insulation.

Applicants respectfully submit that the claims are in condition for allowance, and an early

notification to that effect is earnestly solicited.

Please charge any additional fees or credit any overpayment to Deposit Order Account

No. 11-0855.

Respectfully submitted,

Bruce D. Gray

Reg. No. 35,799

Attorney for Assignee

OF COUNSEL:

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Suite 2800

1100 Peachtree Street

Atlanta, Georgia, 30309-4530

404-815-6218

Attorney Docket No.: M8540/248465

SALINE SOLUBLE INORGANIC FIRRES

This invention relates to saline soluble. non-metallic, amorphous, inorganic oxide, refractory fibrous materials.

Inorganic fibrous materials are well known and widely used for many purposes (e.g. as thermal or acoustic insulation in bulk, mat, or blanket form, as vacuum-formed shapes, as vacuum formed boards and papers, and as ropes, yarns or textiles; as a reinforcing fibre for building materials; as a constituent of brake blocks for vehicles). In most of these applications the properties for which inorganic fibrous materials are used require resistance to heat, and often resistance to aggressive chemical environments.

Inorganic fibrous materials can be either glassy or crystalline. Asbestos is an inorganic fibrous material one form of which has been strongly implicated in respiratory disease.

It is still not clear what the causative mechanism is that relates some asbestos with disease but some researchers believe that the mechanism is mechanical and size related. Asbestos of a critical size can pierce cells in the body and so, through long and repeated cell injury, have a bad effect on health.

whether this mechanism is true or not regulatory agencies have indicated a desire to categorise any inorganic fibre product that has a respiratory fraction as 'hazardous', regardless of whether there is any evidence to support such categorisation. Unfortunately for many of the applications for which inorganic fibres are used, there are no realistic substitutes.

Accordingly there is a demand for inorganic fibres that will pose as little risk as possible (if any) and for which there are objective grounds to believe them safe.

A line of study has proposed that if inorganic fibres were made that were sufficiently soluble in physiological fluids that their residence time in the human body was short; then damage would not occur or at least be minimised. As the risk of asbestos-linked disease appears to depend very much on the length of exposure this idea appears reasonable. Asbestos is extremely insoluble.

As intercellular fluid is saline in nature the importance of fibre solubility in saline solution has long been recognised. If fibres are soluble in physiological saline solution then, provided the dissolved components are not toxic, the fibres should be safer than fibres which are not so soluble. The shorter the time a fibre is resident in the body the less damage it can do. H. Förster in 'The behaviour of mineral fibres in physiological solutions' (Proceedings of 1982 WHO IARC Conference, Copenhagen, Volume 2, pages 27-55(1988)) discussed the behaviour of commercially produced mineral fibres in physiological saline solutions. Fibres of widely varying solubility were discussed.

International Patent Application No. W087/05007 disclosed that fibres comprising magnesia, silica, calcia and less than 10 wt% alumina are soluble in saline solution. The solubilities of the fibres disclosed were in terms of parts per million of silicon (extracted from the silica-containing material of the fibre) present in a saline solution after 5 hours of exposure. The highest value revealed in the examples had a silicon level of 67 ppm. In contrast, and adjusted to the same regime of measurement, the highest level disclosed in the Förster paper was equivalent to approximately 1 ppm. Conversely if the highest value revealed in the International Patent Application was converted to the same measurement

regime as the Förster paper it would have an extraction rate of 901,500 mg Si/kg fibre - i.e. some 69 times higher than any of the fibres Förster tested, and the fibres that had the highest extraction rate in the Förster test were glass fibres which had high alkali contents and so would have a low melting point. This is convincingly better performance even taking into account factors such as differences in test solutions and duration of experiment.

International Patent Application No. WO89/12032 disclosed additional fibres soluble in saline solution and discusses some of the constituents that may be present in such fibres.

European Patent Application No. 0399320 disclosed glass fibres having a high physiological solubility.

Further patent specifications disclosing selection of fibres for their saline solubility are European 0412878 and 0459897, French 2662687 and 2662688, PCT WO86/04807 and WO90/02713.

The refractoriness of the fibres disclosed in these various prior art documents varies considerable. The maximum service temperature of any of these disclosed fibres (when used as refractory insulation) is up to 815°C (1500°F).

Service temperature for refractory insulation is definable in many ways but to be consistent with the above mentioned International Patent Applications this application shall mean by service temperature that temperature at which the fibre shows acceptable shrinkage (maximum of 5% linear shrinkage after exposure to temperature for 24 hours) and at which the fibre has not appreciably suffered through excessive sintering or softening.

There is a demand for physiologically soluble fibres having a service temperature of greater than 815°C,

particularly for such fibres having a service temperature above 900°C.

Testing for physiological solubility and safety can be done by inhalation studies on, e.g. rats. However such studies are extremely time consuming and costly. A study can take of the order of $2\frac{1}{2}$ years from start and can easily cost f1 million per study. A cheaper alternative is to test for solubility in physiological or like fluids in vitro.

Testing of an inorganic fibre for solubility in physiological solutions is not so time consuming, but there is currently no way of predicting which systems will produce such soluble fibres. Therefore anyone seeking to find such soluble fibres has to work on a trial and error basis assisted by what is commonly known as 'chemical intuition' but is equally commonly known as 'a hunch'. Such trial and error testing is laborious and time consuming. Further, once a fibre is found that is soluble there is no guarantee that it will be usable at useful service temperatures.

Accordingly there is a demand for a method of predicting whether a fibre will have a reasonable solubility in physiological solutions, and further there is a demand that such a test should preferably give an indication as to expected service temperature.

Shrinkage of inorganic refractory fibres occurs through two mechanisms; the first is viscous flow of the fibre material. Most inorganic refractory fibres are glasses and so may be defined as liquids having an exceedingly high viscosity (but still liable to flow). By their nature fibres are elongate and so have a high surface area per unit volume. As the reduction of surface area is a means of reducing the surface energy of a material, when the glass becomes fluid enough it will flow so as to reduce surface area. This flow results in a coarsening and shortening of the fibres and so to

shrinkage, and in the extreme results in disruption of the fibres into separate particles.

The second mechanism leading to shrinkage is that at elevated temperatures glasses may crystallise to form one or more crystal phases. Usually these crystal phases have a smaller molar volume than the glasses from which they crystallise and so shrinkage results. Some fibres are known for which the molar volume of the crystalline form exceeds that of the glass (for example ${\rm Al}_2{\rm O}_3/{\rm SiO}_2$ glassy fibres may crystallise to form mullite crystals). In these cases the expansion due to crystallisation may oppose the shrinkage caused by viscous flow.

If shrinkage through viscous flow occurs at a much lower temperature than crystallisation then the crystallisation may not be able to compensate for such shrinkage.

There is a demand for a fibre in which both viscous flow and crystallisation occur at as high and as similar a temperature as possible, and preferably in which the expansion due to crystallisation closely matches the shrinkage due to viscous flow so that the net effect is as close to zero shrinkage as possible.

When used as refractory insulation inorganic refractory fibres are used in several forms. The fibres may be supplied as a bulk material, but in this form the fibres are difficult to handle for many applications. Alternatively the fibre may be supplied as a blanket. Blanket fibre is generally made by a process of sucking fibre from air onto a conveyor to form a blanket. Because the fibres tend to be aligned parallel to the conveyor surface they can separate easily. Accordingly the blanket fibres are secured together by adding a binder to lock the fibres together, or by needling the blanket, or both. In needling needles are passed through the thickness of the blanket to push and draw fibres to lie transverse to the

blanket and so tie the fibres together. Because binders are usually resins, such as phenolic resins, they burn off on first firing. There is a desire to reduce the amount of such binders used both because of possible health implications in handling, and because the combustion products may affect the strength of the fibres. Thus needled blanket is usually preferred.

The fibres may also be supplied as blocks, generally made from assembled layers of inorganic fibre blanket.

For some fibres needling is not possible. Crystalline fibres—are generally too brittle to stand the stresses involved. For the fibres known in the industry as glass fibres (which are generally used for low temperature applications) the amount of 'shot' (unfiberised glass particles) present is generally too high to allow needling as the shot damages the needles. There is no needled blanket on the market that has a maximum service temperature in the range 900°C-1200°C. There are needled blankets having a higher maximum service temperature but these use expensive fibres in comparison with other fibres usable (with the aid of binders) as blanket in the temperature range 900°C-1200°C.

Accordingly there is a demand for needled fibre blanket formed from inexpensive materials, being soluble in saline solutions, and having a maximum service temperature in the range 900°C-1200°C.

As stated previously refractory oxide fibres are made by several methods all of which involve the formation of a melt of oxides and the subsequent fiberisation of the melt by e.g. spinning or blowing.

The melt of oxide material is often formed by electrical discharge melting of the constituent raw materials. The applicants, in manufacture of a CaO/MgO/SiO₂ refractory oxide

fibre encountered problems due to the necessity of handling CaO. These problems were discovered to be due to the moisture content of CaO as commercially available. One of the problems of use of CaO is the outgasing that results upon melting and this led at the least to a porous melt pool which caused fluctuations in the melt current; in the extreme the outgasing was explosive. Additionally use of CaO appeared to cause accelerated attack on the melt electrodes. Also CaO is a difficult and corrosive material to handle.

Accordingly there is a need for a process that minimises the use of CaO.

Accordingly the present invention provides the following features both independently and in combination:-

- A. Use of a vitreous inorganic fibre in the knowledge that it has a composition meeting the criteria either: that the calculated sum of the free energies of hydration of the compounds that would or could be present at equilibrium (on the basis of knowledge, informed belief or reasonable assumption) is more negative than -10 kcal/100grams (-418.4kJ/kg) of composition; or that the percentage of non-bridging oxygens is more than 30%. Such compositions tend to be saline soluble.
- B. Use of such a vitreous inorganic fibre in the knowledge that it has a composition meeting the criterion that the ratio of glass modifiers to network formers is less than a critical value (for SiO₂ based compositions the critical value is 1). Such compositions tend to be glass formers.
 - <u>C</u>. The invention also encompasses fibres selected by adopting such criteria as a test for solubility and glass formation.
 - D. Use as saline soluble fibres having a shrinkage of less

than 3.5% when exposed to 1000°C for 24 hours and/or 800°C for 24 hours, of vitreous fibres having a composition comprising (in weight %):-

$$SiO_2$$
 >58% - (for MgO =< 10%) and SiO_2 >58% + 0.5(%MgO -10) - (for MgO >= 10%) (SiO₂ preferably being below 70%) CaO 0% - 42% MgO 0% - 31.33% Al₂O₃ 0% - <3.97%

and being essentially free of fluxing components such as alkali metals and boron oxide.

 \underline{E} . In one such usage the first crystalline material resulting on crystallisation has the crystal structure of diopside and has the composition consisting essentially of:-

Component	Composition A
	Weight percent
SiO ₂	59-64
Al ₂ O ₃	0-3.5
CaO	19-23
MgO	14-17

F. In a second such usage the first crystalline material resulting on crystallisation has the crystal structure of wollastonite/pseudowollastonite and has the composition consisting essentially of:-

Component	Composition B
	Weight percent
sio ₂	60-67
Al ₂ 0 ₃	0-3.5
CaO	26-35
MgO	4-6

- G. The fibres used in such manner may further be used as needled blankets.
- H. Preferably the fibres of the general composition and compositions A and B mentioned above have a SiO₂ content (expressed as a weight percentage of the constituents SiO₂, CaO and MgO) of greater than 60%.
- I. The present invention further provides a method for the manufacture of refractory oxide fibres containing calcium and silicon by the formation of an oxide melt containing calcium and silicon characterised in that all or part of the calcium and all or part of silicon is provided by a calcium silicate.

The invention is illustrated by way of example in the following description and with reference to the drawings in which:-

- Fig.1 is a three-axis phase diagram indicating the crystalline phases in the system SiO₂/CaO/MgO (*Phase Diagrams* for Ceramists, The American Ceramic Society, 1964) a key to this diagram is at the end of the specification;
- Fig.2 is a three-axis composition plot of the projection onto the $SiO_2/CaO/MgO$ phase field of compositions comprising SiO_2 , CaO, MgO and Al_2O_3 ;
- Fig.3 is a temperature/time plot of the firing regime used in a series of cyclic exposure tests of experimental

compositions;

Fig.4 is a plot of log (total solubility) v calculated free energy of hydration for a series of fibres.

Fig. 5 is a plot of log (total solubility) v % non-bonding oxygens for a series of fibres (see below).

A series of fibres were made of the compositions shown in Table 1. These fibres were melt spun by using a vertical spinning system of the type known for making inorganic fibres. Also shown in Table 1 are the compositions of some comparative commercially available inorganic oxide fibres and glass fibres.

TABLE 1

	Al ₂ 0 ₃	SiO ₂	CaO	MgO	zro ₂	
SW-A SW-A1 SW-A2	3.3 1.1 0.8	59.3 6 × 63.7 6 60.8 6 *	20.5	15.5 15.2 15.4	-	
SW-B1 SW-B2 SW-B3	2.3 1.3 1.0	65.3 66.9 60.0	26.8 27.5 34.0	5.7 5.2 4.4	- -	
	A1 ₂ 0 ₃	sio ₂	COMPARA CaO	ATIVE H	EXAMPLES ZrO ₂	MST
CRBT CWBT CHBT	46.5 40.6 49.7	53 49.5 35.1	0.04 5.50 0.04	0.01 4.00 0.01	- - 14.7	1260°C 870°C 1425°C
Glass Fibre	15.2/ 15.5	53.7/ 57.5	21.1/21.8	1.3/		+ 5.9-6.2% B ₂ O ₃ 0.11-0.12% TiO ₂ 0.46% Na ₂ O 0.32-0.33% K ₂ O
Needled Glass Fibre	3.7	60.5/ 60.0	8.1 7.9	4.0		+ 2.85-2.95% B ₂ O ₃ 13.5% Na ₂ O 1.0% K ₂ O

[MST = Maximum Service Temperature (oxidising atmosphere)]

The fibres SW-A, SW-A1, SW-A2, SW-B1, SW-B2 and SW-B3 were tested for solubility by the following method.

The fibre was first chopped in the following manner. 2.5 g of fibre (deshotted by hand) was liquidised with 250 cm 3 of distilled water in a domestic Moulinex (Trade Mark) food blender for 20 seconds. The suspension was then transferred to a 500 cm 3 plastic beaker and allowed to settle after which as much liquid as possible was decanted and the remaining liquid removed by drying in an oven at 110°C.

The solubility test apparatus comprised a shaking incubator water bath, and the test solution had the following composition:-

Compound	<u>Name</u>	<u>Grams</u>
NaCl	Sodium chloride	6.780
NH ₄ Cl	Ammonium chloride	0.540
NaHCO ₃	Sodium bicarbonate	2.270
$\text{Na}_2\text{HPO}_4.\text{H}_2\text{O}$	Disodium hydrogen	0.170
	phosphate	
Na ₃ C ₆ H ₅ O _{7P} .2H ₂ O	Sodium citrate	0.060
	dihydrate	
H2NCH2CO2H	Glycine	0.450
H_2SO_4 s.g. 1.84	Sulphuric acid	0.050

The above materials were diluted to 1 litre with distilled water to form a physiological-like saline solution.

0.500 grams \pm 0.0003 grams of chopped fibre was weighed into a plastic centrifuge tube and 25 cm³ of the above saline solution added. The fibre and saline solution was shaken well and inserted into the shaking incubator water bath maintained at body temperature (37°C \pm 1°C). The shaker speed was set at 20 cycles/minute.

After the desired period (usually 5 hours or 24 hours)

the centrifuge tube was removed and centrifuged at ≈ 4500 revs/minute for approximately 5 minutes. Supernatant liquid was then drawn off using a syringe and hypodermic needle. The needle was then removed from the syringe, air expelled from the syringe, and the liquid passed through a filter (0.45 micron cellulose nitrate membrane filter paper [WCN type from Whatman Labsales Limited]) into a clean plastic bottle. The liquid was then analysed by atomic absorption using a Thermo Jarrell Ash Smith - Hiefje II machine.

The operating conditions were:-

ELEMENT	WAVELENGTH(nm)	BAND WIDTH	CURRENT (MA)	FLAME (Nitrous Oxide + Acetylene)
Al	309.3	1.0	8	Fuel Rich
${\tt SiO}_2$	251.6	0.3	12	16 11
CaO	422.7	1.0	7	Fuel Lean
MgO	285.2	1.0	3	f1 11

The procedure and standards adopted for determining the above elements were as set out below.

 SiO_2 can be determined without dilution up to 250 ppm concentration (1 ppm = 1mg/Litre). Above this concentration an appropriate dilution was made volumetrically. A 0.1% KCl solution (0.1g in 100 cm³) was added to the final dilution to prevent ionic interference. NB If glass apparatus is used, prompt analysis is necessary.

From a stock solution of 1000 ppm pure ignited silica (99.999%) (fused with Na_2CO_3 at $1200\,^{\circ}C$ for 20 minutes in a platinum crucible (0.2500g $SiO_2/2g$ Na_2CO_3) and dissolved in dilute hydrochloric acid (4 molar) made up to $250\,\mathrm{cm}^3$ with distilled water in a plastic volumetric flask) the following

standards were produced:-

STANDARD (PPM SiO ₂)	STOCK SOLUTION (cm ³)
10.0	1.0
20.0	2.0
30.0	3.0
50.0	5.0
100.0	10.0
250.0	25.0

Add 0.1% KC1 to each standard before making to 100cm3.

Aluminium may be measured directly from the sample without dilution. Standards of 1.0, 5.0 and 10.0 ppm Al may be used. For calibration readings are multiplied by 1.8895 to convert from Al to Al_2O_3 .

A standard Al atomic absorption solution (e.g. BDH 1000 ppm Al) was bought and diluted using an accurate pipette to the desired concentration. 0.1% KCl was added to prevent ionic interference.

Calcium may require dilutions on the sample before determination can be carried out (i.e. \times 10 and \times 20 dilutions). Dilutions must contain 0.1% KC1.

A standard Ca atomic absorption solution (e.g. BDH 1000 ppm Ca) was diluted with distilled water and an accurate pipette to give standards of 0.5, 4.0 and 10.0 ppm. 0.1% KC1 is added to prevent ionic interference. To convert readings obtained from Ca to CaO a factor of 1.4 was used.

Magnesium may require dilutions on the sample before determinations can be made (i.e. \times 10 and \times 20). Add 0.1% KC1 to each dilution. To convert Mg to MgO multiply by 1.658.

A standard Mg atomic absorption solution (e.g. BDH 1000 ppm Mg) was diluted with distilled water and an accurate

pipette to give standards of 0.5, 1.0 and 10.0 ppm Mg. 0.1% KC1 was added to prevent ionic interference.

All stock solutions were stored in plastic bottles.

The results of the tests are indicated in Table 2.

TABLE 2

BODY FLUIDS SOLUBILITY

(ppm)

		SiO ₂	Ca	aO	. 1	lg0
	5 h	24h	5h	24h	5h	24h
SW-A	98	120	63	56	33	66
SW-A1	83	141	32	70	21	70
SW-A2	130	202	43	73	100	177
SW-B1	58	77	10	38	5	9
SW-B2	64	121	27	55	5	10
SW-B3	138	192	80	46	8	21

Fibres with the best solubility (SW-A2 and SW-B3) were then tested, after annealing at varying temperatures, and compared with the comparative examples of Table 1. The results are shown in Table 3.

It can be seen that for the SW-A2 fibre, with increasing annealing temperature, the silica solubility drops progressively. In contrast the SW-B3 composition shows no loss in solubility up to 800°C and although a reduction in solubility is shown above that temperature it is not as dramatic as for SW-A2. Despite this difference in solubility it is to be noted that only the needled GF fibre shows a

comparable silica solubility and that material melts at 700°C.

TABLE 3

<u>Fibre</u>	Condition	S	olubili	ty Ana	lyses		
		CaO	(ppm)	MgO	(ppm)	Sio	(ppm)
		5hrs	24hrs	5hrs	24hrs	-	24hrs
SW-A2	As received	58	37	37	3	8 9	130
SW-A2	600°C, 48hrs	33	56	27	43	60	108
SW-A2	800°C, 48hrs	35	53	17	30	43	87
SW-A2	1000°C, 48hrs	7	3	3	2	11	21
SW-B3	As received	35	6 9	7	22	22	100
	600°C, 48hrs			12	22	55	
SW-B3	800°C, 48hrs				7	24	
SW-B3	1000°C, 48hrs			3	3		60
CRBT	As received	10	8	6	3	5	3
CHBT	As received	16	10	7	3	4	0.3
Glass Fib	re As received	14	17	5	3	5	7
	F As received		34	8	15	6 6	85
Needled G	F 600°C, 48 hrs	11	26	7	10	19	37
Mineral F	ibre As received	16	16	7	6	8	9

[The Glass Fibre and Needled Glass Fibre had the compositions shown in Table 1.]

The user is primarily concerned with the solubility of the fibre as received as it is in this condition that most handling occurs; as received both SW-A2 and SW-B3 fibres have extremely high solubility. Even after exposure to 800°C and 1000°C these fibres have solubilities much higher than other high temperature use fibres.

To investigate the reasons underlying the difference in solubilities after high temperature annealing between the SW-A2 and SW-B3 fibres qualitative X-ray diffraction was done on the fibres. The results are indicated in Table 4 and it can be seen that the SW-B3 fibre forms pseudowollastonite and wollastonite, whereas the SW-A2 fibre forms diopside. It appears therefore that the crystalline diopside has a lower solubility in physiological saline solution than the crystalline pseudowollastonite and wollastonite material precipitated from the SW-B3 fibre.

TABLE 4

Sample	Condition	Qualitative XRD
SW-A2 SW-A2	600°C, 48 hours 800°C, 48 hours	Amorphous Amorphous with small amount
SW-A2	1000°C, 48 hours	Diopside Diopside
SW-B3	600°C, 48 hours	Amorphous
SW-B3	800°C, 48 hours	Amorphous
SW-B3	1000°C, 48 hours	Pseudowollastonite & Wollastonite

Various of the fibres were then tested for their shrinkage characteristics. Table 5 shows the results of Shrinkage tests on all the test fibres and on some of the comparative fibres. These results were obtained by proposed ISO standard ISO/TC33/SC2/N220 (equivalent to British Standard

BS 1920, part 6,1986) with some modifications to account for small sample size. The method in summary comprises the manufacture of vacuum cast preforms, using 75g of fibre in 500cm³ of 0.2% starch solution, into a 120 x 65mm tool. Platinum pins (approx 0.1-0.3mm diameter) were placed 100 x 45mm apart in the 4 corners. The longest lengths (L1 & L2) and the diagonals (L3 & L4) were measured to an accuracy of ±0.01mm using a travelling microscope attached to a steel rule with a vernier scale. The samples were placed in a furnace at temperature and left for 24 hours. The shrinkage values are given as an average of the 4 measurements.

TABLE 5
LINEAR SHRINKAGE (%)

(24h at temperature)

Temp.°C	SW-A	SW-Al	SW-A2	SW-B1	SW-B2	SW-B3
730	1.45		1.43	1.02	0.22	
870				0.41	·	
900			1.07			1.07
1000		1.04	1.3	0.51	0.6	1.1
1100		0.71	1.8		0.73	2.2
Maximum Service Temperatur °C	850 e	1050	1050	1050	1050	1000

It can be seen that in SW-A, SW-A1, SW-A2, SW-B1, SW-B2 and SW-B3, owing to the increase in molar volume on crystallisation, the linear shrinkage at the maximum service temperature is less than 3.5%.

Table 6 shows the results of a further series of shrinkage tests made in the same way.

TABLE 6

Sample	Measurement Direction c.f. Roll	Test Temperatures °C	Linear Shrinkages % Range Mean
	Direction		
<i></i>			
SW-A2	Parallel	850	1.1-1.4 1.2
SW-A2	Perpendicular	850	0.7-1.5 1.3
SW-A2	Parallel	900	0.5-1.1 0.9
SW-A2	Perpendicular	900	1.9-4.5 3.0
SW-A2	Parallel	1000	0.5-2.9 1.3
SW-A2	Perpendicular	1000	1.7-2.9 2.2
SW-A2	Parallel	1100	0.7-1.5 1.0
SW-A2	Perpendicular	1100	1.0-2.6 1.8
SW-B3	Parallel	900	1.6-1.8 1.7
SW-B3	Perpendicular	900	1.4-2.4 2.1
SW-B3	Parallel	1000	1.6-2.3 1.9
SW-B3	Perpendicular	1000	1.0-2.3 1.7
SW-B3	Parallel and	1100	Complete Meltin
	Perpendicular		(Lantern type remnant)

To ascertain the applicability of these tests to long term usage a series of cyclic shrinkage tests were undertaken on the materials and the heating schedule used for these cyclic

tests is shown in Figure 3.

The results of the tests are shown in Tables 7 & 8 (the two figures given for SW-B3 are due to slight differences in chemical analysis [the fibre at the end of a production run of fibre tends to have slightly differing composition to that at the beginning of a production run of fibre]).

As a further comparison with the above discussed materials a melt was made comprising 55% SiO₂, 29.9% CaO and 18.6% MgO. Fibres made using this composition had a maximum service temperature of 700°C and melted at 800°C.

As these results were encouraging the applicants conducted a further and extensive series of tests, concentrating on the SW-A2 and SW-B3 compositions, to ascertain the reproducibility of these results and the boundaries of the useful compositions.

Table 9 (three pages) below gives the compositions of a series of melts, ranked on silica content, and showing the shrinkage figure after exposure to 1000°C for 24 hours (1st column) and 800°C for 24 hours (2nd column). shrinkages were measured by the same method as the shrinkages given above but measurements were made with a travelling microscope with a digital linear scale accurate to \pm 5 μ m. can clearly be seen that all fibres with a silica content of less than 58% have a shrinkage at 1000°C of greater than 3.5% save two (B3-3 and 708). These fibres, together with some fibres with a silica content of greater than 58% although showing a reasonable figure at 1000°C, show a very poor figure at 800°C. Compositions with an SiO2 content of greater than 70% appear to fiberise poorly. This may be because such compositions have two liquids in the melt as appreciated from Fig.1

TABLE 7

CYCLIC SHRINKAGE (LINEAR)

20

(왕)

Product	1000°C	1100°C	24h at 1000°C
No. cycles	58	42	
CRBT	2.0	2.7	1.9
CWBT	15.0	13.3	12.1
SW-A2	0.33	2.0	1.3
SW-B3	1.00	1.67	1.1
SW-B3	0.33	0.67	1.1

Accuracy: + or - 0.33%

TABLE 8
CYCLIC SHRINKAGE

(%)

-		LINEAR SHRINKAGE		THICK SHRIN	
PRODUCT	1000°C	1100°C	24 h at 1000°C	1000°C	1100°C
No. cycles	104	100		104	100
CRBT CWBT SW-A2 SW-B3 SW-B3	1.47 14.4 1.5 1.73 1.47	3.1 15.2 2.1 1.63 1.77	1.9 12.1 1.3 1.1	0.47 38.63 8.58 7.24 7.02	11.19 32.14 8.75 7.57 7.16

Accuracy: + or - 0.3%(%)

TABI.R. C

	%	9	5	8	7	7	8	8	7	c	0	4	9	7	8	7	c	8	2	7	2	8	6	4	9	_	8	7	0		æ
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Shrinka at °C	1000	-	0.7			•	1.1	•	•	•	•		•			•	•	•	•	•	•	•	•	•		•				1.4	
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	F																													7	
itions	K20%	0.0	0	0.0	0	0.0	•	0.0	0.0	٠.	•	0.0	•	0.		۰.	•	•	•	•		0	Ţ.	0	0.	0	0	0	0	0.13	0
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	ca0%	0.	4.	.3	4.	2.6	د .	4.	. 7	.7	1.5	4.6	3.2	5.1	9	8.1	0.9	7.4	5	8.4	7.7	4.9	9.9	4.8	1.2	6.2	9.8	5.6	8.7	25.81	6.8
	sio_2 %	8.0	3.4	3.0	2,3	2.2	1.4	1.2	1.1	0.8	0.4	0.0	9.4	9.2	8.7	8.6	8.5	8.3	8.1	7.6	7.5	7.5	7.2	9.9	6.1	6.1	5.8	5.7	5.6	55.50	5.3
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	Me1	2-	2-	2-	3-	2-	2	2	2	3-	2	2-	3	2-	3.	2	3-	59	2	2	7	3	3	2	5	2-	3-	19	N	B3-	N

TABLE 9 (continued)

Analysed Compositio (Weight %) 38 Al ₂ O ₃ & Na ₂ O ₈ K ₂ 54 0.58 0.43 0.05 53 0.06 0.17 <0 58 0.34 0.17 <0 59 0.34 0.17 <0 50 0.29 0.10 <0 50 0.29 0.10 <0 50 0.29 0.10 <0 50 0.29 0.10 <0 50 0.20 0.30 0.11 <0 50 0.20 0.27 0.27 <0 60 0.25 <0 60 0.25 <0 60 0.25 <0 60 0.25 <0 60 0.25 <0 60 0.26 0.36 0.37 <0 60 0.27 <0 60 0.25 <0 60 0.26 0.36 0.36 <0 60 0.27 <0 60 0.27 <0 60 0.28 0.38 0.38 0.38 <0 60 0.28 0.38 0.38 0.38 <0 60 0.28 0.38 0.38 0.38 0.38 <0 60 0.25 0.39 0.30 0.30 0.30 0.30 0.30 0.30 0.30	ns Shrinkage Rationalised at °C Compositions	0% Fe ₂ O ₃ % ZrO ₂ % 1000 800 SiO ₂ % CaO% MgO%	7	05 0.20 0.49 0.4 65.7 27.3 7.	9 0.19 0.61 1.0 68.1 26.1 5.	5 0.08 <0.05 0.3 0.6 66.6 27.9 5.	5 0.16 0.80 3.4 3.9 66.4 6.8 26	5 0.11 0.13 3.2 3.3 65.6 14.0 20	9 0.14 0.28 0.6 66.5 33.1	05 0.13 0.47 7.3 7.1 65.5 9.1 25	09 0.16 0.30 1.0 66.3 24.0 9	12 0.17 0.58 7.4 4.3	07 0.15 <0.05 2.7 1.6 65.0 16.4 18	10 0.11 <0.05 1.9 0.9 64.9 21.9 13.	05 0.08 0.05 1,0 2.3 64	11 0.15 0.15 21.6 18.8	$05 \mid 0.13 \mid <0.05 \mid 0.4 \mid 0.5 \mid 64.0 \mid 30.4 \mid 5.$	05 0.16 <0.05 1.3 2.9 63.4 21.2 15.	05 0.16 0.07 6.1 63.9 11.3 24.	09 0.17 0.91 0.8 0.9 65.6 32.2 2.	05 0.10 0.23 1.4 63.7 24.0 12.	35 0.17 <0.05 0.6 63.4 28.9 7.	12 0.25 0.59 0.9 63.4 34.2 2.	10 0.17 <0.05 1.8 1.2 63.6 26.4 10.	05 0.14 0.08 3.3 12.	05 0.16 <0.05 1.3 62.8 31.8 5.	15 0.09 0.66 0.3 3.0 64.1 20.7 15.	15 0.19 0.12 7.5 3.	05 0.13 0.16 1.	10 0.15 0.09 2.5 1.7	0.09 0.08 1.8 1.1 61.7 24.8 13.	0.28 0.49 1.6 1.7 62.0 35.4 2	0.15 0.05 18.9 10.7 61.7 19.1 19.
102 & Ca0 & La O & Cao & La O & Cao	lysed (We	90% A1 ₂ O ₃ % Na	06 /0 06	60.07 66.	.54 0.58	5.33 0.06	$6.20 \mid 0.23 \mid 0$	9.98 0.34	0.37 0.64	4.88 0.29	.33 0.56	.68 3.97 0	.21 0.40 0	.96 1.49 0	.00 0.33 0	.59 5.70 0	.44 <0.05 0	.22 0.20 <0	.18 0.33 0	.06 0.55 0	.98 0.44 0	.54 0.10 0	.33 0.69 0	.73 0.58 0	74 0.31 0	25 0.05 0	54 2.57 <0	48 4.15 0	.66 0.28	.99 1.74	.24 0.25	50 0.76	78 0.48
		io28 cao8	5 23 27 14	3.63 67.14 5.11 3.4 6.14	20, 22, 21, 26, 27	2.72 80.6	4.85 6.63	4.16 13.74 1	4.13 31.93	4.12 8.88	4.09 23.26	3.74 25.41	3.68 16.06 1	3.66 21.44 1	3.56 16.55 1	3.24 24.83	2.61 29.79	2.60 20.92 1	2.36 10.99 2	2.33 30.62	1.98 23.37 1	1.83 28.13	1.71 33.25	1.62 25.53	1.38 13.62 2	1.33 31.08	1.32 19.78 1	0.83 32.30	0.74 25.30 1	0.32 32.27	0.32 24.28 1	0.28 34.49	0.20 18.59 1

TABLE 9 (continued)

			Ana]	Analysed Co (Weight	mpos	itions			Shrinkaç at °C	ıkage °c	Rationa Composi	nali siti	sed
Melt	$\sin_2 k$	CaO&	%06W	A12038	Na208	K208	Fe2038	Zro2%	1000	800	$\sin_2 x$	Ca0%	Mg0%
~	0.1	2.5	8	.7	Ŋ	.1	. 2	8	•		2	<u>س</u>	
932	59.85	21.60	15.65	1.50	90.0	<0.0>	0.18	0.11	12.8		61.6	22.3	16.1
92	9.8	4.3	4.	٦.	۲.	•	0		•		0	4.	2
3-3	9.5	1.6	9.	Q.	۳.	<u>.</u>	7	4.	•	15.3			
Ċ	9.5	8.1	7	4.	.3	•	۲.	0.	•		-	9	•
3-2	9.4	0.9	9	9	4.		7	۲.	•		1.	7.	٠
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734	6.5	9	•	•	0	•	۲.	0			58.2	24.3	17.5
7	6.1	4.4	6.8	J.	0	<0.05	Ţ	0	4.		7.	5	7.
(C)	5.9	0.9	4.	Š	4.	•	د .	4	•	34.7	ъ В	7.	•
0	5.2	2.7		•	3	<0.05	4	0	Ή.	•	<u>ي</u>	.	0
71	4.6	4.0	9	Ñ	<u>_</u>	<0.05	7	<0.0>	_		i,	4.	•
										7			

*Poor fibres containing a lot of shot. All other constituents < 0.1%. Too poor to test for solubility or shrinkage

PARIE 10

90% A1203 & Na20% K20% Fe203 & ZrO2 & SiO2 CaO MgO SiO2 & CaO				Anal	ysed (Weig	Composit pt %)	ions		Ť	Solubil ppm	ilit opm	ies	Ratio Compo	ionalis positi	sed
2-28 78.07 17.15 0.15 <0.05	elt	10_2	ao	g	$^{1}_{2}^{0}_{3}$	a ₂ 0	20	e203	ro_2	0.1	ਰ	מ ו	10_2	Ø	Mg0%
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2-28	3.0	2.0	7.1	۲.	0.0	0.0	۲.	.7	F			0	۲.	7.
9. 5.3 0.9 6.36 19.60 0.23 0.05 <0.05	2-16	3.4	12.4	0.0	7	0.0	0.0	۲.	.2	\leftarrow			9	5	•
1-32 72.38 23.43 0.65 0.31 0.09 0.22 0.72 4 15.0 35.48 75.0 24.25 12.35 111 <0.05 <0.05 0.10 0.13 1.03 35.48 74.3 13.3	2-32	3.0	6.3	9.6	3	•	0.0	7		-			.	6.	6
2-15 72.25 12.67 12.35 0.11 <0.05	3-32	2.3	23.4	•	4	٠.	0.		.7				5.	4.	•
2-2 71.48 9.36 16.34 0.33 0.10 <0.05	2-15	2.2	12.6	2.3	7	0.	0.0	٦,		0			4	э •	2.
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2-22	1.4	9.3	6.3	۳.	۲.	0.0	?	8	2		7	ب	•	9
2-27 71.14 6.77 19.64 0.24 0.08 <0.05	2-29	1.2	4.4	2.3	7	0.	0.0		.2	9		0	5	•	•
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2-27	1.1	6.7	9.6	7	0.	0.0	•	•	0			5	7.	0
2-17 70.43 11.58 14.52 0.15 <0.05	3-28	9.8	18.7	0.	4.	.2	0.	-	.7	0			ب	9.	•
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2-17	7.4	11.5	4.5		0.	0.0		5	9			н Н	2	•
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2-33	0.0	4.6	2.8	.2	Ţ	0.0		0				٠.	4.	<u>ب</u>
2-6 69.29 15.17 12.76 0.07 0.25 <0.05 0.11 1.13 152 48 70 71.3 15.0 3-16 68.74 24.99 1.71 0.65 0.38 0.10 0.24 1.03 132 72 10 72.0 26.2 2-26 68.65 8.12 19.26 0.29 <0.05	3-18	9.4	23.2	3.7	4.	۳.	0.		4.	4			;	4.	رب
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	9-2	9.2	15.1	2.7	0	.2	0.		۲.	Ω			1.	5.	٠
2-26 68.65 8.12 19.26 0.29 <0.05 <0.05 0.21 0.98 183 31 97 71.5 8.3 33-27 68.56 20.98 7.00 0.78 0.35 0.08 0.17 0.79 122 54 20 71.0 21.0 2-20 68.13 17.45 12.60 <0.05	3-16	3.7	24.9	1.7	9	.3	۲.		0.	3			8	9	1:
59 68.56 20.98 7.00 0.78 0.35 0.08 0.17 0.79 122 54 20 71.0 21.0 29 68.33 17.45 12.60 <0.05	2-26	9.6	8.1	9.5	3	0.	0	7	6.	8			;	ж Э	•
59 68.33 17.45 12.60 <0.05 0.33 <0.05 117 38 40 69.5 17. 2-20 68.19 11.58 16.57 0.40 0.19 0.05 0.21 0.91 162 31 69 70.8 12. 2-31 67.62 8.45 21.72 0.32 0.09 <0.05	3-27	8.5	20.9	7.0	.7	. 3	0.		.7	2			;	.	7.
2-20 68.19 11.58 16.57 0.40 0.19 0.05 0.21 0.91 162 31 69 70.8 12. 2-31 67.62 8.45 21.72 0.32 0.09 <0.05	59	8.3	17.4	2.6	0	۲.	0	٠,	0	_			6	7.	5
2-31 67.62 8.45 21.72 0.32 0.09 <0.05	2-20	8.1	11.5	6.5	4.		0.	?	9	9			0	5	٠
3-31 67.59 27.76 0.49 0.40 0.11 0.22 1.01 112 50 4 70.5 29. 3-19 67.58 24.91 3.65 0.45 0.37 0.07 0.15 0.65 163 44 21 70.3 25. 3-17 67.25 26.68 1.86 0.70 0.45 0.10 0.23 0.57 162 49 13 70.2 27. 2-13 66.67 14.87 16.01 0.11 0.05 <0.05	2-31	7.6	8.4	1.7	۳.	0.	0.	-	•	0			6	α •	22.2
3-19 67.58 24.91 3.65 0.45 0.37 0.07 0.15 0.65 162 49 13 70.3 25.5 3-17 67.25 26.68 1.86 0.70 0.45 0.10 0.23 0.57 162 49 13 70.2 27. 2-13 66.67 14.87 16.01 0.11 0.05 <0.05	3-31	7.5	27.7	4.	4.	4.	7		0				0	ნ	٠
3-17 67.25 26.68 1.86 0.70 0.45 0.10 0.23 0.57 162 49 13 70.2 27. 2-13 66.67 14.87 16.01 0.11 0.05 <0.05	3-19	7.5	24.9	•	4.	۳.	0		•	9			0	س	٠
2-13 66.67 14.87 16.01 0.11 0.05 <0.05 0.10 0.92 159 47 70 68.4 15. 2-10 66.17 16.22 15.80 0.49 0.06 <0.05 0.09 <0.05 122 42 52 67.4 16. 3-22 66.17 21.28 9.34 0.52 0.33 0.08 0.18 0.54 174 62 57 68.4 22. 3-15 65.86 29.82 1.78 0.45 0.05 0.03 <0.51 156 38 7 67.6 30. 19 65.77 25.69 8.12 <0.05 0.24 <0.05 0.23 <0.05 115 51 24 66.0 25. 2-5 65.69 18.74 13.78 0.18 0.16 <0.05 0.10 0.14 150 48 60 66.9 19. 3-4A 65.50 25.81 4.88 2.05 0.28 0.13 <0.05 0.14 11.7 181 54 84 67.7 17.	3-17	7.2	26.6	ω.		4.	Ξ.	7	ശ	9			0	7:	;
2-10 66.17 16.22 15.80 0.49 0.06 <0.05 0.09 <0.05 122 42 52 67.4 16.33-2 66.17 21.28 9.34 0.52 0.33 0.08 0.18 0.54 174 62 57 68.4 22.3-15 65.86 29.82 1.78 0.47 0.45 0.08 0.18 0.51 156 38 7 67.6 30.19 65.77 25.69 8.12 <0.05 0.24 <0.05 0.23 <0.05 115 51 24 66.0 25.2 25.2 65.69 18.74 13.78 0.18 0.16 <0.05 0.10 0.14 150 48 60 66.9 19.3 3.44 65.50 25.81 4.88 2.05 0.22 0.13 <0.05 0.14 11.7 181 54 84 67.7 17.	2-13	9.9	14.8	6.0	۲.	•	•	7	o.	വ			æ	٠ د	
3-22 66.17 21.28 9.34 0.52 0.33 0.08 0.18 0.54 174 62 57 68.4 22. 3-15 65.86 29.82 1.78 0.47 0.45 0.08 0.18 0.51 156 38 7 67.6 30. 19 65.77 25.69 8.12 <0.05 0.24 <0.05 0.23 <0.05 115 51 24 66.0 25. 2-5 65.69 18.74 13.78 0.18 0.16 <0.05 0.10 0.14 150 48 60 66.9 19. 3-4A 65.50 25.81 4.88 2.05 0.28 0.12 0.16 0.24 82 52 16 2-8 65.33 16.86 14.24 0.22 0.13 <0.05 0.14 1.17 181 54 84 67.7 17.	2-10	6.1	16.2	5.8	4.	•	0	0	0	2				9	•
3-15 65.86 29.82 1.78 0.47 0.45 0.08 0.18 0.51 156 38 7 67.6 30. 19 65.77 25.69 8.12 <0.05 0.24 <0.05 0.23 <0.05 115 51 24 66.0 25. 2-5 65.69 18.74 13.78 0.18 0.16 <0.05 0.10 0.14 150 48 60 66.9 19. 3-4A 65.50 25.81 4.88 2.05 0.28 0.12 0.16 0.24 82 52 16 2-8 65.33 16.86 14.24 0.22 0.13 <0.05 0.14 1.17 181 54 84 67.7 17.	3-22	6.1	21.2	٠,	r.		0.	;	.5	/			8	S	•
19 65.77 25.69 8.12 <0.05 0.24 <0.05 0.23 <0.05 115 51 24 66.0 25. 2-5 65.69 18.74 13.78 0.18 0.16 <0.05 0.10 0.14 150 48 60 66.9 19. 3-4A 65.50 25.81 4.88 2.05 0.28 0.12 0.16 0.24 82 52 16 2-8 65.33 16.86 14.24 0.22 0.13 <0.05 0.14 1.17 181 54 84 67.7 17.	3-15	5.8	29.8		4.	4.	0.	۲.	.5	Ω			7	0	•
2-5 65.69 18.74 13.78 0.18 0.16 <0.05 0.10 0.14 150 48 60 66.9 19. 3-4A 65.50 25.81 4.88 2.05 0.28 0.12 0.16 0.24 82 52 16 2-8 65.33 16.86 14.24 0.22 0.13 <0.05 0.14 1.17 181 54 84 67.7 17.	19	5.7	25.6	۲.	0.	?	0.	. 2	0.	_			9	2	ω
3-4A 65.50 25.81 4.88 2.05 0.28 0.12 0.16 0.24 82 52 16 2.8 2.8 65.33 16.86 14.24 0.22 0.13 <0.05 0.14 1.17 181 54 84 67.7 17.	2-5	5.6	18.7				0.	۲.	7	S			9	٠ ق	•
2-8 65,33 16,86 14,24 0,22 0,13 <0.05 0,14 1,17 181 54 84 67,7 17.	3-4A	ນ	25.8	8	0.			۲.	7	8					
	2-8	5.3	16.8	?	7	<u>:</u>	0.	.	۲.	8			7	/	14.8

TABLE 10 (continued)

			Ana]	lysed Co (Weight	Composit ht %)	tions		,	Solubili ppm	ilit pm	ies	Rationa Composi	ali iti	sed
Melt 8	${\rm Sio}_2 \$$	ca0%	MgO&	A12038	$\mathrm{Na}_2\mathrm{O}\$$	K208	Fe2038	Zro28	$\sin 2$	cao	Мдо	$\sin 2 $	Ca0%	Mg0%
18	5.2	7.1	6.	0	.2	0.	~	4.	0	47	20	5.	7.	•
3-14	5.1	9	ß	٠ د	4.	0	. 1	9.	158	67	27	æ	•	•
21	5.0	7.2	۳.	0.	7	0.0	0	0.	S	89	20	9	7.	•
2-34	4.8	9.9	6.2	.2	•	•	-	æ.	3	11	119	9	6.	9
2-21	4.1	3.7	ę.	.3		0.0	۲.	·ı	9	28	56	5.	4.	•
3-30	4.	31.93	ö	0.64	0.45	0.09	0.14	0.28	163	83	m	66.5	33.1	0.4
2-35	4.1	8.8	ω.	.2		0.	۲.	4.	9	18	106	5	9.	•
3-23	4.0	3.2	9.3	ა.	٠,	0.	7	.	101	44	31	9	•	٠
3-5A	3.7	4.	9.	6.	7.	۲.		.5	4	43	11			
2-30	3.6	6.0	8.2	4.		0.	۲.	•	0	33	52	5	6.	α
5-9	3.6	1.4	2.9	4.	٠,	۲.	~	0.	9	67	99	4	21.9	13.2
2-12	3.5	6.5	0	е		•	0	•	9	47	99	•	•	8
3-6A	3.2	4.8	4.5	. 7		۲.	٠,		2	7	26			
23	2.6	9.7	4.	0.		0.		•	4	62	17	4.	0	•
57	2.6	0.9	5.2	~	•	0	۲.	0.0	Ø	62	7	ب	;	•
2-25	2.3	0.9	۲.	۳.		•	٦.	•	9	35	105		;	4.
3-13	2.3	9.0	2.0	ς.	•	0.	.	6.	S	41	7	<u>ي</u>	5	7
2-7	1.9	3.3	و	4.		o.	۲.	?	7	29	63	ب	4.	•
25	1.8	8.1	7.5	4	٠,	0.	٦.	•	186	92	35	63.4	28.9	7.7
3-11	1.7	3.2	۳.	•	ა.	۲.	7	ິນ	9	28	15	ж •	4.	<u>ن</u>
3-24	1.6	5.5		٠ ر	۳.	۲.	٠,	•	œ	37	57	ن	9	•
2-24	1.3	3.6	.,	.	7	0.	Ţ.	•	m	27	87	2	ر	ر
22	1.3	1.0	5.2	0.	7	0	۲.	•	8	81	23	۲.	;	Ω.
24	1.3	9.7	3	3	۰.	0.	0.	9.	7	36	37	4.	0	•
3-2A	0.8	2.3	0.4		٤,	۲.	۲.	۲.	S	61				
2-14	0.7	5.3	9	7	۲.	0.			127	57	43	62.2	25.9	11.9
3-1A	0.3	2.2	3.9	.7	٠,	۲.	٦.	0.	~	72				
2-11	0.3	4.2	2	.2		0.	0.	0	4	54		;	4.	•
3-9	0.2	4.4	2.5	. 7	3		7.	4.	2			62.0	ك	2
23	60.20	8.5	.7	4.		0.	۲.	0.	g	35	47	-	19.1	19.5
								4		1	7		1	1

TABLE 10 (continued)

Rationalised Compositions	Mg0%		16.1	5		•	•	•	•	•		6.2	•	•		•		•	8	•	7	7			- 1
	Ca0%	۳.	22.3	4.		6	7.	N.	ς.	6	ល	33.9	4.	δ.	ж	6.	æ	i.	7.	4.	4.	5	7.	•	4
	Sio ₂ %	2	61.6	0		-	;	9.	1.	0	0	59.9	9.	9	9	6	9	8	6		8	7	ъ В	٠ کا	
ies	оБу	17	43	23	10	33	ည	30	37	99	19	27	169	23	40	22	16	41	57	73	99	47	18	7	62
bilit ppm	cao	84	41	67	25	48	29	73	48	42	54	99	16	69	29	24	25	64	47	55	52	42	14	31	55
Solubilit ppm	\sin_2	146	92	140	30	133	0	4	9	~	8	133	0	2	8	9	2	4	9	4	\sim	0	8	9	c
	ZrO_2	8.	۲.	-	4.	<0.05		0	.2	0	0		0.	0.	0.	0.	0.	0.	0.	•	0.	0	4.	0.	<0.05
	Fe2038	.2		0	~	0.14	7	4.	•		+		.2	۲.	7			•	۲.			۲.	۳.	₽.	.2
ions	K20%		0	0.	~	0.08	۲.	•	0	0	0	0.	0.	0.		0.	۲.		۲.	•	•	0		0	
Analysed Compositions (Weight %)	Na20%	5	0.	۲.	۳.	0.39	4.	2	۲.	3	7	. 2	0.	7	ß	3	ن	4.	5	۲.	0	0	4.	۵.	۲.
	A12038	.7		۲.	6	0.48	ę.	0.	7.	3		7.	7		ō.	0.	.7	•	9	.7	0	Š	5	0.	.2
	Mg0%	8	9	4.	•	•	•	.2	~	.7	9	•	0.	æ	0	•	4.	4.	3.0	٤.	7.0	6.8	4.4	.7	19.66
	Ca0%	2.5	1.6	4.3	1.6	8	6.0	2.0	0.9	8.4	5.0	3.1	3.9	4.6	2.7	5,3	9.9	0.5	6.2	رن ق	3.6	4.4	6.0	2.7	
	Sio ₂ %	0.1	9.8	9.8	9.5	9	9.4	0.6	8.8	8.7	8.6	8.3	7.7	7.7	7.5	7.5	7.3	6.9	6.9	6.8	6.5	6.1	5.9	5.2	54.68
	Melt	C	\sim	9	3-3	B3-25	3 - 2	14	9	~	8	9	9	9	C	12	3-2	3	2-1	7	C	7	C	08	71

There are several anomalies, namely compositions B3-6A, A2-25, A2-24, A2-23, B3-2A, B3-3A, A2-19, and 932. All of these have an SiO_2 content of > 58% but a high shrinkage.

On the assumption that the minimum silica level for satisfactory shrinkage varies with MgO content the applicants have determined that fibres with a silica content (in weight percent) that fail to meet the following expression do not have satisfactory shrinkages at either or both 800°C and 1000°C:-

$$SiO_2$$
 >58% - (for MgO =< 10%) and SiO_2 >58% + 0.5(%MgO -10) - (for MgO >= 10%)

The applicants have further found that the ${\rm Al}_2{\rm O}_3$ content is important. From their studies it appears that the maximum ${\rm Al}_2{\rm O}_3$ content lies somewhere between 2.57% and 3.97%. The applicants have found that with increasing alumina levels the first material to crystallise is calcium aluminate and this possibly forms a liquid phase that assists flow and hence shrinkage.

Table 10 shows, for the same compositions as Table 9, 24 hour solubilities for each major constituent. It can be seen that all of the compositions have high solubilities.

As mentioned above use of CaO in forming calcium containing fibres is inconvenient and can be hazardous. The applicants investigated use of mixed oxide materials that would avoid the handling of CaO. A fibre was made by admixture of magnesia with silica and wollastonite (CaSiO₃).

The raw materials used to make the melt comprised:-

Pennine Darlington Heavy Magnesia (#200)

MgO	-	92.60%
CaO	-	1.75%
Fe_2O_3	-	0.04%
sio ₂	-	0.20%
Cl	-	0.25%
so ₃	-	0.70%
LOI	-	4.50%

Partek's Finnish Low Iron Wollastonite (#200) (U.K. agent - Cornelius Chemical Co., Romford, Essex)

sio_2	-	51.80%
CaO	-	44.50%
MgO	-	0.80%
A1 ₂ 0 ₃	-	0.60%
Fe_2O_3	-	0.30%
Na ₂ O	_	0.10%
к ₂ 0	-	0.05%
TiO2	-	0.05%
S	-	0.02%
MnO	-	0.01%
P	-	0.01%
F	-	0.01%
LOI	-	1.70%

Hepworth Mineral's Redhill T washed silica sand

SiO₂ - 99.0% min.

These constituents were mixed as 78.65% Wollastonite; 19.25% SiO₂; and 3.6% MgO. This gave 0.4 - 0.5% of the final melt as Al_2O_3 .

It was surprisingly found that in producing a melt using these constituents the current requirements were only two-thirds that for the use of the raw oxides.

Fibre was produced by blowing (although spinning and other methods can be used). 2 runs were performed with different blowing conditions.

Chemical analysis was undertaken by the Analytical Department at the applicant's subsidiary Morgan Materials Technology (hereinafter called ${\tt M}^2{\tt T}$) using wet chemical techniques. Fibre diameters were measured using ${\tt M}^2{\tt T}$'s Galai particle analyser, with shape analysis software. Typically 40,000 fibres were analysed for each run.

The first result of note was the speed of melt reaction when using wollastonite as compared with lime. Also the current was seen to be very stable throughout the growth of the melt. If the current was lost whilst pulling the electrodes apart the current could be restored simply by pushing them back together again. This was not possible with the runs using lime.

Chemical Analysis

	<u>Others</u>	<u>CaO</u>	MgO	Al ₂ 0 ₃	<u>sio</u> 2	zro ₂	Na ₂ O	Total
Run1	0.7	32.6	3.8	0.8	60.1	0.8	0.5	99.3
Run2	0.7	32.5	3.8	0.8	60.1	0.8	0.6	99.3

Runs 1 and 2 indicate the respective x-ray fluorescence analyses for each run.

Shrinkage Results (1000°C for 24 hours)

		<u>L1</u>	<u>L2</u>	<u>L3</u>	<u>L4</u>	Av.	Std.Dev.
Run	1	0.9	0.2	0.4	0.6	0.5	0.3
Run	2(A)	1.0	-0.2	0.7	0.6	0.5	0.5
Run	2(B)	0.5	0.2	0.0	0.4	0.2	0.2

Solubility Results (ppm)

	<u>CaO</u>	<u>Mg0</u>	SiO ₂
Run 1(5 hr)	67	10	95
Run 1(24hr)	84	17	146
Run 2(5hr)	39	7	72
Run 2(24hr)	73	17	186

Fibre diameters

	<u>Mean</u>	Median	100%<	<u>%>5µm</u>	%<1µm
Run 1	5.1 μ m	3.4 μ m	30μm	33%	13%
Run 2	$4.1 \mu \mathrm{m}$	$2.7 \mu m$	25µm	25%	19%

Accordingly it appears to be the case that by using what are cheaper ingredients than the pure oxides one can obtain a fibre that has as high a performance as using purer oxides and at much improved energy costs and safety. It is to be noted that this feature of the invention is not limited to saline soluble fibres and any oxide fibre that contains both calcium and silicon can advantageously be made with a calcium silicate, wollastonite being merely an example of such a silicate.

The previous description is directed towards high temperature usage of particular saline soluble fibres. The following is directed towards the prediction and use of saline soluble fibres. A series of fritted glass melts were made of the compositions shown in Tables 11A and 11B and quenched in water. Solubilities of the various components of the quenched melt were measured by the previously described method of atomic absorption. The solubilities were normalised to a specific surface area of 0.25m²/gram of fibre.

The free energy of hydration was calculated by normalising the chemical analysis to 100 weight %; making the assumption that simple silicates (MSiO $_3/M_2$ SiO $_3$) are present

and calculating the free energy of hydration contribution of each species; and summing to get the total free energy of hydration. The data in Tables 11A & 11B is also presented in Fig.4. It can be seen that the fibres lie on a generally straight line bar four groups of materials that will be explained below.

Table 11 shows in each column the following:-

Fibre reference Composition

Molar ratios

Moles/100 grams of melt

Species assumed (see below)

Calculated free energy of hydration of assumed species (kcal/100 grams)(see below)

Calculated free energy of hydration of assumed species (kJ/kg) (see below)

Solubility data (# indicates not measured [see below])

Specific surface area

Normalised solubility data

log normalised solubility

The base data on which calculation of the free energy of hydration was done is set out in Tables 12 which indicates free energies of hydration taken from the literature in the units kcal/mol and kJ/mol.

lable of free Energy of Hydration Values for Silicate Melts

TABLE 11

lable of free tnergy of Hydration Values for Silicate Helts

TABLE 11 Continued

																		_		_		_) j	},															
		solub)						85.L				İ	2.38					1.72				9.30				2 08					70 1	?			1.15	:			
	Not Buil 1 2 ed	Solubiller	28	,				ŝ					212					70			7.6	3		_		119	:				92	<u> </u>			71	:			
		2/3/					3,	0.3v					e. 54				-	20.00			07.0	;				0.48					0.40				0.36				
	<u> </u>		~ ·	·		• \$	8	2 '		- ;	× :	2	<u> </u>	ā		3 :		? -	- «	2	=	. `	~	•	\$	2	25	•	118	228	107	_	2	147	-	0	_	=	20
100	5	3000	3 6	20014	6000	70.00	100		054	AL 203	2016	1000	2 4	20014	2018		-	2001	\$102	Total	Q.J	Cox	A1203	\$102	lotal	CaO	Мво	A1203	2015	lotal	810	A1205	2015	Total	K20	MgO	A1 203	2015	Total
free freenon of	Hotension of the state	(ga/ca) 100 .			181.0	¥ 097-	7 607	C HE	3.66	7. 5	1 105.	1 702	180 4		8.10	0 277	7.607		-102.1	7.987	197.5	29.6	55.4	-177.6	-349.3	-260.8	-226.4	1.0	59.0	-437.2	-421.7	22.9	-101.4	-500.2	-305.4	-188.0	22.9	204.8	-265.7
free Energy of	Hydration (kgal /100a)	.7.2	-0.2	0.7	.4.3	0.11.	-9.8		2.0	7.6	-12.5	.7.8	17.	0.0	1.5	10.6	8.6.	9.0	-2.4	-11.6	1.4.	1.0.	7.7	-4.2	.a.3	-6.2	-5.4	0.0	1.4	-10.2	1.01.	0.5	-2.4	-12.0	-7.3	.2.1	0.5	2.1	-6.B
Holes	in 1kg	2.96	0.30	2.39	2.50		4.01	1.22	87.1	1.38		3.18	3.10	0.10	2.64		10.4	1.93	1.41		2.93	1.09	4 .26	3.24		3.87	3.89	0.08	2.52		4.13	1.76	1.40		1.73	3.23	1.76	8.75	
Species		Srsio3	Q,H	A1203	Sro		Srsio3	35	A1203	Sro		Srsios	Musio3	A1203	2015		Sisios	A1203	810		Casios	H _U C)	A1205	CaO		CaSi03	MgS iO3	AI 203	S i 02		Srsio3	A1203	SrO		K25103	MgSiO3	A1203	2015	
Moles	in 100g	0.296	0.030	0.239	0.250		0.401	0.122	0.146	0.138		0.318	0.310	0.010	992.0		0.401	0.193	0.141		0.293	601.0	0.426	928.0		0.387	0.389	0.008	0.252		0.413	0.176	0.140		0.175	0.323	0.176	0.377	
Holes		0.546	0.030	0.239	9.530		0.539	0.122	0.148	0.401		0.318	0.310	0.010	0.892		0.542	0.193	0.401		0.617	O. 109	0.426	0.293		0.387	0.389	0.00	1.026		0.553	0.176	0.413		0.175	0.323	0.176	0.675	
	Rationalized	56.6	1.2	7.72	17.8	100.0	55.9	6.7	15.1	24.1	100.0	32.9	12.5	1.0	53.6	100.0	2.95	19.7	24.1	0.001	34.6	4.4	43.4	17.6	100.0	21.7	15.7	\$2 '	61.6	100.0	57.3	17.9	24.8	100.0	16.5	13.0	17.9	.52.6	100.0
Composition		55.0	1.2	23.7	17.5	97.2	55.0	4.8	2.0	23.7	98.3	33.0	12.5	9.	53.8	156.3	26.0	19.7	24.0	2.66	35.0	5.7	0.33	2 8	101.3	21.7	15.7	8.3	9.10	0.00	\$6.5	17.6	54.4	98.5	16.5	2.0	16.0	52.B	100.3
CO	oxide oxide	Sr0	N ₉ 0	A1203	2015	lotal	Sro	O. H	A1203	2013	lutal	Sr0	M ₉ 0	A1203	2018	letal	Sro	A1205	2015	lotal	Ca0	0f.W	A1203	2015	1013	ÇaQ	g.	A1203	2015	letol	Sro	A1203	S i 02	lotal	K20	₩ 06 ₩	A1203	Sio2	Total
Metr Code		SMAS(A)					SMAS(B)					SHS(A)					SAS(C)				CMAS(A)					SMA2					SAS(D)				KHAS(A)				

TABLE 12

			G_{hyd}	
Releva	ant oxides		(kcal/mole)	(kJ/mol)
_	+ H ₂ O> eous silica)	H_2Sio_3	5.6	23.4
		231(OH)	3.1	12 0
	+ 3H ₂ O>	-		13.0
	+ H ₂ O>	-	- 6.5	-27.2
	+ H ₂ O>	_		- 54.8
	+ H ₂ O>	_	-17.3	- 72.4
-	+ H ₂ O>		-33.5	-140.2
_	+ H ₂ O>		-46.1	-192.9
_	+ H ₂ O>	-		66.9
	+ 3H ₂ O>		-55.9	-233.9
	+ 3H ₂ O>		- 9.8	-41.0
Zro ₂	+ H ₂ O>	Zr(OH) ₂ O	-7.1	-29.7
Relevant Si	<u>ilicates</u>			
Na ₂ SiO ₃ +2H	+(aq)> H ₂ S	iO ₃ +2Na ⁺ (aq)	-28.8	-120.5
	$^{L}(aq) \longrightarrow H_2S$	-		-174.5
	⁺ (aq)> H ₂ S			-58.2
	(aq)> H ₂ S			-67.4
	+(aq)> H ₂ S			-102.1
-	$^{+}(aq) \longrightarrow H_2S$		-37.3	-156.1
•	+(aq)> H ₂ S	•	-2.4	-10.0
,	2	J		
Disilicates	<u>5</u>			
Ca ₂ SiO ₄ +3H	+(aq)> H ₂ S	iO ₃ +2Ca ²⁺ +OH ⁻	-30.3	-126.8
	(aq)> H ₂ S			-210.5
	$^{\dagger}(aq) \longrightarrow H_2S$			- 74.9
		3	- · ·	

Although calculations were undertaken assuming the presence of the simplest silicates choice of other silicates (such as disilicates) does not appear to change the calculations much. For example given below is the calculation for an assumed composition which shows only minor differences for the free energy of hydration calculated.

```
0.767 CaO
Composition (moles/100g)
                                     0 417 MgO
                                     0.669 SiO2
0.384 \text{ Ca}_2\text{SiO}_4 =
                        -11.6
                                     or
                                           0.433 CaSiO<sub>3</sub>
                                                                    -7.0
0.285 MgSiO3
                                           0.236 MgSiO3
                         - 4.0
                                                                    -3.3
0.132 MgO
                         - 0.9
                                           0.334 CaO
                                                                    -4.4
                         -16.5
                                           0.181 MgO
                                                                    -1.2
                                                                   -15.9
                                          0.384 \text{ Ca}_{2}\text{SiO}_{4} =
                                     or
                                                                    -11.6
                                            0.132 \text{ Mg}_2 \text{SiO}_4 =
                                                                     - 2.4
                                            0.153 MgSiO<sub>3</sub>
                                                                     <u>-2.1</u>
                                                                     -16.1
```

The applicants have found that when the free energy of hydration is more negative than -10kcal/100 grams (-418.4kJ/kg) of composition the composition showed high solubility. The compositions where this relationship broke down were those for which the total solubility was not available (for example those materials containing sodium, where any dissolved sodium would be swamped by the sodium in the saline solution) or where the free energy of hydration of the most likely species present was not available from the literature.

As a test of this technique the two examples of European Patent No. 0399320 were examined. The disclosed examples had the compositions:-

Component	Composition 1	Composition 2
	Weight percent	Weight percent
sio ₂	60.7	58.5
Al ₂ O ₃	-	5.8
CaO	16.5	3.0
Mg0	3.2	-
B ₂ O ₃	3.3	11.0
Na ₂ O	15.4	9.8
к ₂ 0	0.7	2.9
Iron oxide	0.2	0.1
BaO	-	5.0
ZnO	-	3.9

Using the above method of calculation Composition 1 had a free energy of hydration of -11.6kcal/100grams (-485.3kJ/kg) whereas Composition 2 had a free energy of hydration of -5.8kcal/100grams (-242.6kJ/kg). This would suggest that Composition 1 would be a saline soluble fibre, and hence physiologically safer than an insoluble fibre; whereas Composition 2 would be predicted to be a relatively insoluble fibre and hence less safe. This is what is disclosed in EP 0399320, the fibres of Composition 2 having a longer lifetime introduced fibres were which the studies in in interperitoneally into rats.

As mentioned above this predictive test can fail under some circumstances. To avoid these difficulties the applicants looked to a different predictive technique, namely the assessment of the amount of non-bridging oxygens present. This is calculated by normalising the chemical analysis to 100 weight%; calculating the molar percentage of each oxide; summing the oxygen-weighted contribution of each oxide to get the total number of oxygens; summing the weighted contribution of each oxide of non-bridging oxygens (see below); and taking the ratio of non-bridging oxygens to the total number of oxygens. The applicants have found that when this figure exceeds 30% the fibres are soluble.

To explain the term non-bridging oxygen one must look to the structure of glasses. Glasses are very stiff liquids and to form usually require the presence of a material that can form a network (usually an oxygen-bridged network). The network may be modified by constituents that contribute non-bridging parts to the network and open the structure of the network and so prevent crystallisation. These materials are usually referred to as network-formers and modifiers respectively.

The terms modifier and network former are well known in the glass industries. Network formers are materials such as SiO_2 , P_2O_5 , B_2O_3 and GeO_2 which can form an interconnected network to form the glassy phase. Modifiers are substances such as CaO, Na_2O , and K_2O which alter the network and have effects on such properties as viscosity and melting point. There are some intermediate materials (such as Al_2O_3 , TiO_2 , PbO, ZnO and BeO) which can act as both network formers and modifiers depending on the environment and on the amount present.

In the above mentioned test, for calculating the non-bridging oxygens, one ignores the network formers and calculates the contribution of each other oxide. The contribution of each oxide depends on the geometry and charge of each cation in the glass. As examples typical contributions are as follows:-

Ca²⁺,Mg²⁺,Sr²⁺ and other divalent network modifier cations contribute 2 non-bridging oxygens

K⁺,Na⁺ and other monovalent network modifier cations contribute 1 non-bridging oxygen

 ${\rm Al}^{3+}, {\rm Ti}^{3+}$ and other intermediate cations contribute -1 non-bridging oxygen (i.e these oxides <u>reduce</u> the number of non-bridging oxygens)

 $(\mathrm{Ti}^{4+}\ \mathrm{is}\ \mathrm{reduced}\ \mathrm{to}\ \mathrm{Ti}^{3+}\ \mathrm{in}\ \mathrm{most}\ \mathrm{glasses}\ \mathrm{when}\ \mathrm{present}\ \mathrm{in}\ \mathrm{relatively}\ \mathrm{small}\ \mathrm{quantities})$

38
TABLE 13

Composition Holes
0.842
0.240
27.1 0.451 29.4 fotal 1.533 100.0
0.829
0.232
0.473
1.534 100.0
0.852
0.234
0.451
1.537
_
7.7 (80.0)
1.602
09.0
34.9 0.581 48.8
0.00
40.1 0.022 40.0
1.692
0.303
0.389
1000 49.7 1 100 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
0.374
0.347
44.2 0.736 50.5
+
0.331
0.882
1.461
0.305
0.612
1.205
0.338
0.374
43.6 0.726 50.5
1

TABLE13 (continued)

	2		Hotes	Mol.X	Oxygen	.0.g.	XN.B.0	Solub	Solubility	4 0 0	1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1	
	e le	אנ א			lotat			Oxide	(Dxxm)	27.70	Day I I Feb.	L og(norm
SHAS(A)	ors O	55.0	0.531	49.1	169.6	8.09	35.8	Sro	25	27.0	ATT TOTAL	solub)
	Š.	1.2		2.8				CoM	, v	5	8	1.58
	A1203	23.7	0.232	21.5				A1203	۰ ۰			
	2015	17.3		56.6				Si02				
		lotal	1.081	100.0				Total	, %			
SHAS(B)	Sro	55.0		44.6	157.6	84.8	53.8	Sro	110	02.0	,	
	M ₉ 0	8.7	0.119	10.0				o Go	ž	6.39	Ş	1.98
	A1203	14.8	0.145	12.2				A1203	<u> </u>			
	Sio2	23.7	0.394	33.2				Cois	- 2			
		Total	1.189	100.0				2016	77.			
SHS(A)	SrO	33.0	0.318	20.7	159.5	80.6	\$ 05	000	140	Š		
	MgO	12.5		20.2		2	}	0 9	<u>}</u> ;	0.39	212	2.33
	A1203	1.0	0.010	9.0				2001	ē •			
	Si02	53.8	0.895	58.4				6 6013	o ĉ			
		Total	1.533	100.0				Total	Ç.			
SAS(C)	Sro	56.0	0.540	47.7	169.3	61.4	36.3	Sro	5	02.0		
	A1203	19.7	0.193	17.0				¥1204	? -	97.78	7	1.72
	Sio2	24.0	0.399	35.3				Sio	- «			
		Total	1.132	100.0				lotat	۶ ۾			
CHAS(A)	CaO	35.0		42.6	179.3	41.4	23.1	CaO		02.0	1,6	
	N ₉ 0	4.5	0.112	7.6				Ob H		0.00	3	1.58
	A1203	0.77	•	29.5				A1203	M			
	Si02	17.8	0.296	20.3				\$102	000			
		lotal	1.464	100.0				Total	82			
SWAZ	Q :	21.7		21.4	157.5	85.0	54.0	CaO	53	0.48	110	2 OB
	<u> </u>	15.7	0.389	21.5				MgO	25		:	2
	A1203	8.0		9.0				A1203				
	2015	61.8	1.028	2.95				Si02	118			
		Total	1.812	100.0				Total	228			
SAS(D)	Szo	56.5	0.545	48.5	166.9	5.99	39.7	Sro	107	07 0	60	100
	A1203	17.6	0.173	15.4				A1203	-	?	7,	<u>.</u>
	Si02	54.4		36.1				5102	30			
		Total	1.124	100.0				Total	147			
KHAS(A)	8	16.5		11.3	179.5	41.4	23.1	K20	3	0.36	71	1 15
	M ₉₀	13.0		20.8				06#	. 00	3	:	?
	A1203	18.0	0.177	11.4				A1203	. –			
	Sio2	52.8		56.6				S i 02	-			
		Total	755	0 001								

Table 13 shows, for the compositions of Table 11, the calculated figure for non-bonding oxygens and these figures are reproduced in Fig.5. It can be seen that the plot of Fig.5 is more linear than that of Fig.4. Compositions with greater than 30% non-bridging oxygens show high solubility.

To illustrate the method the calculation for one example (the first shown in Table 13) is given below:-

	Mols/	Total Oxygen	Non-bridging Oxygens
CaO Al ₂ O ₃ SiO ₂	0.549 0.157 0.294	0.549 0.471 (0.157*3) 0.588	1.098 (2*0.549) -0.314 (-1*0.157*2Al) 0.000
	1.000	1.608	0.784

% non-bridging oxygens (0.784/1.608)*100 = 48.8%

As an example one can look to European Patent Specification No. 0399320 referred to above. Using this method in relation to that specification Composition 1 has a non-bridging oxygen percentage of 48.2% whereas Composition 2 has a non-bridging oxygen percentage of 19.6%, again predicting that Composition 1 is more soluble than Composition 2.

There is a further criterion which a composition must meet for it to form vitreous fibres, namely that it be capable of forming a glass. The applicants have found a simple test. If the ratio of modifiers to network formers is less than a critical value (for SiO₂ based glasses, 1) the composition will generally form a glass. For the purpose of this test reasonable results are obtained if such intermediate materials are treated as networkers. Table 14 shows for a series of compositions in each column:-

Fibre reference (N.B. these are not the same fibres as shown in tables 9 & 10)

Composition

Molar ratios

Ratio of glass modifiers to network formers

Free energy of hydration of raw oxides

Melting Point

X-ray diffraction results

Solubility data (# indicates not measured)

Specific surface area

Normalised solubility data

Arbitrary ranking as to solubility and glass forming ability

Indication as to whether melting point above 1500°C

It should be emphasised that this test is a screening one rather than a wholly predictive one as there are several circumstances that may lead to its failure. Among these circumstances are compound formation and inability to quench fast enough to form glass.

Having adopted these tests as a screening method there follows a further step to ascertain whether the composition will form a vitreous fibre. This last step is best examined experimentally as fibre forming ability is a complex function of many physical characteristics, e.g. viscosity, which are often difficult to measure.

Key to Fig.1

CR Cristobalite

TR Tridymite

PS Pseudowollastonite

WO Wollastonite

RA Rankinite

LI Lime

PE Periclase

FO Forsterite

PR Protoenstatite

DI Diopside

AK Akermanite

ME Merwinite

MO Monticellite

TABLE 14

JH. PT	JOSL'S O	•			-	7		1	2	-				-							-			-						F
- IU Rating)	C1858				-						7			-	1		•	1		2		•			2		=		3	100
Criteria (U-10 Rating) H	Sol 45		,	7	•	1				#	•	:			1			Ē		a									~	7
Norm Solub(U.75 m2/9)	Asys		=			1/2		2	SINZ	170	12,				-	50		12			77	•	The second second		1		7		3	WC.
			ř.	112		1.1		ر ا	344	753	13	1.	150	•		316		123		Ĩ	71			•	Ē		~		2	77
	(A)		×.	-0.57		10.53		3.	. O.B.	7,0.71	. u B3	2.5	73.0		1.15	8		ř		J.U.	0.51	_	20.0		22:0		19.0		79.0	75:0_
I E SEL		1	<u> </u>	1_	ž =∽	8	.		22/	27 - Sus	362	31	<u> </u>	?≈	٦	1	375	<u> </u>	~ 	<u> </u>		50		30	ł	×	10,		~	9
III)	<u> </u>	-	909		<u>-</u>	100		*	12			İ	22				333	L	~~			2.		9:		_ =	99	ı	9	
ata tin	22	- 1	20.0		۵ د د	~ -~	20		7221	7921	<u> </u>	k -312	- -	ê E	ř	51	Š	-	200	<u> </u>	-	20		9 2n		8 2	ľ			
<u></u>	• !	اً		1			~		~	61	182	<u> </u>	200		1,4	-	25		252	<u>[</u> ,					<u></u>				*= *	
Soluti	0.00	2005	200	D60	s i 02	41205 A1205	2015	205	510	0.5	000		C30	490 500 500	9	222	C 20 S i 0 2	520	20i2	1000	Reg	A1205	202	A1208		A1205 Si02	Red	2102	A1205	0.55
ARD Data	Americanous Toliaie	American de la companya del companya del companya de la companya d	(No peaks)	Ameng phous . Trace	te santi pents)	(Glassy trace)	CPCREATITION	. :	3	Amiliated Titles	Crystattine (strong)	r ug	(Mod. cryst.)	(No glass)	Crystattine	Crystatthe	some glass (Strong cryst.)	Mediations 1	some crystal. (8 small peaks)	rystaltine	Amoi phous	(No peaks)	Amortidades	(No peaks)	Amor prious	(No peaks)	Amoi phous		(No peaks)	Amengalous Time
		_ K	E .	0531	3	(fut.)	1565		3	550	Š	Ξ	1573		1,565	Ñ		K		3	75	(Eut.)	JIS ST		0551	_	0521	,	ê E	14.00
(kcel/mol (C)			1.3 cf at. 1	-	•	.6.3	1.32.		_		200	17.2	-	·4.3 (fut.)	5:6		. 9 . /	37 1 4		,,,,,,		7.	•	.2.7 E		<u>-</u>	7.6			3.6
Metworker	76.0	0.8		0.2		0.6	25.				16:0	#. D	2		.u/s:>_	1:0		7.7.		1.45	13:0		0.31		0.20		72.D			16.0
\$0	1 0 ALZOS	2.5 C.5	1.0 1 io2 4.5 \$102	<u>~ o</u>	20150	0 A1 205	× 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	1.0 P205	1.0 \$102		. 1 S 202	==	<u> </u>	2015 7 5 107	<u></u>	=	2.1.5:02	- ^	:~i	> 0 > =	-	~ <	- -		Ξ	-~	~~	6 9 5102	1.0 A1205	500
3	223	205 7 23	43 6 S102	5.4	62 4 Si02	~~	53:3 CM	50.6 6205	15.9 Si02	57.2 5102	57.5 Mg0 62.5 S102	C1.7 C10		2015	75.5 C30-	-028 W 3 02	20:50	15 3 600	42.7 \$102	48.2 6205	21 0 11320	50.1 A1205	023 0 02	30.4 A1205	21.2 tid	44.6 \$102	15.9 Red 15.9 A 203	-	32.4 A1203	27.5 5.0
- Total	40. 10	CISIAJ	2	MAS(A) Mo. 20	TAIS.	40. 2I	CPICAL	28: S	No. 23	No. 24	% . %	2317	CHSCAL	,,	Ko. 25	HCS(A)	: \$\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	78.5	*0. 30	No. 31	1	Mpt.	KASIET	£ .	CASK		HASIB) Ho. 35	111313	No. 36	SCS(A)

heleicike

43 14 (continued) TABLE | Hoin | Solub | 10.75 m2/9/1 | Criteria | 10.10 Rating | H. Pt. | 1037 | Crist | Crist | 1500C Ę 507 TEL 3 ż 0.56 U. 8. 0.7 27.0 19:0 _0.62 U. 57. J. 52 חישנ 0.63 10191 256 579 ŏ 369 249 \$ 144 36 Ξ 22 425 = 3 ž 2 7,7,7 3 322 Ş 260 3 22 A1203 S102 S102 Sr0 S102 Sig A1205 C130 C130 C130 C130 C130 A1203 C130 A1203 C130 A1203 lrace (1 smill heat) (Eut) | 1500 CI y8 CB (TING TS CF GIG) | 1501 Amoi proks | (Mo peaks) (fut) some cryst. (neak) then thous to the (1500 C) (3 small peaks) (2 peaks) CTPSTBTTING (No peaks) (No peaks) Anoi Hous (No peaks) Anot phous RRO DSIS ISTER 1 (feat.) S ISSUE US60 C) CJ 995L 100 CO 15 350 Cf.uc) 1600 1121 (Eut.) (17) (17) Cheal/mol (C) 1593 ((111) (E141) -€ ò. S .3.2 7.0 1.00 6.3 3 U.W. U.7B 0.7 37.0 10.0 U.Y 1.05 D. 52 U.Y Ę. 9.0 J.S.D ACT STAT KRAS(A) SCS(C) No. 38 SCS(B) SHAS(8)-SMAS(A) THAS(A) SAS(8) No. 40 SHS(A) SASTET 54 10. 47 5828 10. 48 HAS(C) -5A5(0) -KRASLAF No. 51

CLAIMS

1. A method of predicting whether a given composition will form a saline soluble vitreous material, the method comprising the steps of:-

EITHER

- a) assuming the composition to be a mixture of the compounds that would be present at equilibrium (on the basis of knowledge, informed belief or reasonable assumption);
- b) calculating the sum of the free energies of hydration of each of the compounds assumed to be present;
- c) if the sum of the free energies of hydration is more negative than -10 kcal/100grams (-418.4kJ/kg) of composition, assuming the composition to be soluble in saline solution;

OR

- a") calculating the percentage of non-bridging oxygens present in relation to the total oxygen present and, if the percentage is greater than 30% assuming the composition to be soluble in saline solution;
- 2. A method as claimed in claim 1 which further comprises in combination with steps a)-c) and/or a"):-
- d) calculating the ratio of glass modifier to network former present;
- e) if the ratio is less than a critical value (for SiO₂ based compositions, 1) assuming that the composition will form a vitreous material.

- 3. A saline soluble vitreous inorganic fibre selected and used in the knowledge that it has a composition meeting the criteria that:-
- a) the calculated sum of the free energies of hydration of the compounds that would or could be present at equilibrium (on the basis of knowledge, informed belief or reasonable assumption) is more negative than -10 kcal/100 grams (-418.4kJ/kg) of composition;

and/or

- b) the percentage of non-bridging oxygens present in relation to the total oxygen present is greater than 30%.
- 4. A saline soluble vitreous inorganic fibre as claimed in claim 3 selected and used in the knowledge that it has a composition meeting the criterion that the ratio of glass modifiers to network formers is less than a critical value (for SiO₂ based compositions, 1).
- 5. Use of a saline soluble vitreous inorganic fibre in the knowledge that it has a composition meeting the criteria that:-
- a) the calculated sum of the free energies of hydration of the compounds that would or could be present at equilibrium on the basis of knowledge, informed belief or reasonable assumption is more negative than -10 kcal/100grams (-418.4kJ/kg) of composition;

and/or

- b) the calculated percentage of non-bridging oxygens in relation to the total oxygen content is more than 30%.
- 6. Use as claimed in claim 5 of a saline soluble vitreous inorganic fibre in the knowledge that it has a composition meeting the criterion that the ratio of glass modifiers to network formers is less than a critical value (for SiO₂ based compositions, 1).

7. Use as saline soluble fibres having a shrinkage of less than 3.5% when exposed to 1000°C for 24 hours and/or 800°C for 24 hours, of vitreous fibres having a composition comprising (in weight %):-

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5i0_2 >58% - (for MgO =< 10%) and 5i0_2 >58% + 0.5(%MgO -10) - (for MgO >= 10%) CaO 0% - 42% MgO 0% - 31.33% Al<sub>2</sub>O<sub>3</sub> 0% - <3.97%
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and being essentially free of fluxing components such as alkali metals and boron oxide.

- 8. Use as claimed in claim 7 of saline soluble fibres having a shrinkage of less than 3.5% when exposed to 1000°C and/or 800°C for 24 hours, of vitreous fibres in which the amount of SiO₂ is <70%.
- 9. Use as claimed in any of claims 7-8 of saline soluble fibres having a shrinkage of less than 3.5% when exposed to 1000°C and/or 800°C for 24 hours, the fibres crystallising as diopside and having the composition consisting essentially of:-

Component	Composition A
	Weight percent
SiO ₂	59-64
Al ₂ O ₃	0-3.5
CaO	19-23
MgO	14-17

10. Use as claimed in any of claims 7-8 of saline soluble fibres having a shrinkage of less than 3.5% when exposed to 1000°C and/or 800°C for 24 hours, the fibres crystallising as wollastonite and/or pseudowollastonite and having the composition consisting essentially of:-

Component	Composition B
	Weight percent
SiO ₂	60-67
Al ₂ O ₃	0-3.5
CaO	26-35
MgO	4-6

- 11. Use as a saline soluble fibre as claimed in any of claims 5-10 and in which, owing to the increase in molar volume on crystallisation, the linear shrinkage at the maximum service temperature is less than 3.5%.
- 12. A method for the manufacture of refractory oxide fibres containing calcium and silicon by the formation of an oxide melt containing calcium and silicon characterised in that all or part of the calcium and all or part of silicon is provided by a calcium silicate.
- 13. A method as claimed in claim 13 in which the calcium silicate is wollastonite.

ABSTRACT

Disclosed is use of a vitreous inorganic fibre in the knowledge that it has a composition meeting the criterion that the calculated sum of the free energies of hydration of the compounds that would or could be present at equilibrium (on the basis of knowledge, informed belief or reasonable assumption) is more negative than -10 kcal/100grams of composition. Such compositions are saline soluble.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE



Applicants:

Gary Anthony Jubb and Jean-Louis Martin

Serial No.:

Divisional of 09/262,378

Filing Date:

October 24, 2000

For:

SALINE SOLUBLE INORGANIC FIBRES

Box Patent Application

Date: October 24, 2000

Director of Patents and Trademarks

Washington, D.C. 20231

TRANSMITTAL OF FORMAL DRAWINGS

Sir:

Applicants transmit herewith three sheets of Formal Drawings in connection with the filing of this divisional application. Applicants paid the issue fee in parent U.S. Serial No. 09/262,378 on September 29, 2000, and at that time provided the U.S. Patent Office with identical Formal Drawings.

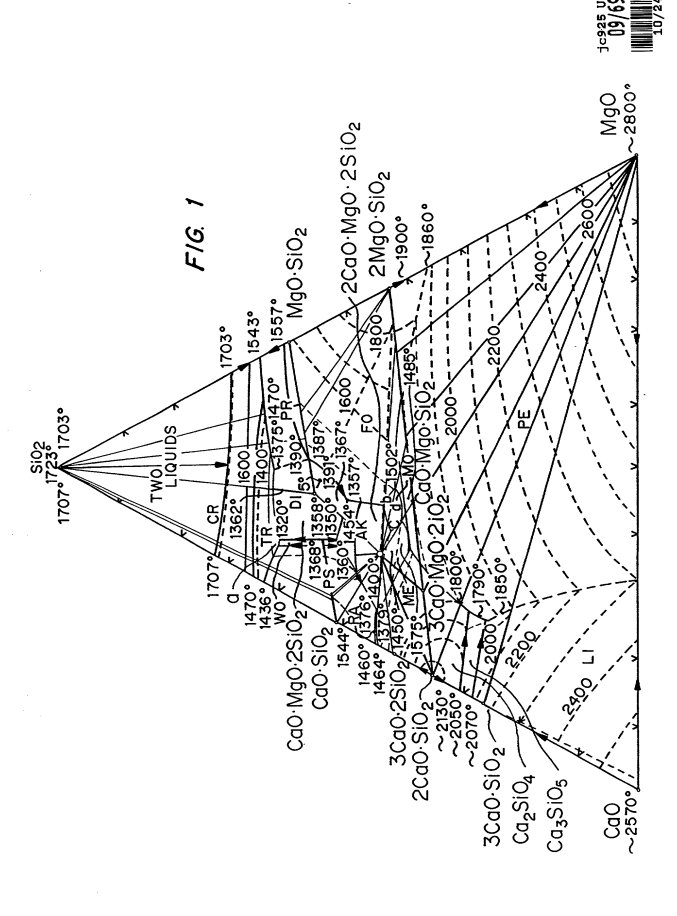
Respectfully submitted,

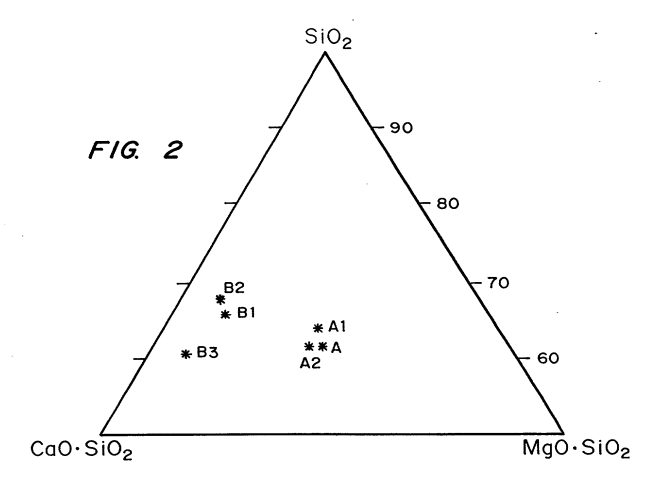
Bruce D. Gray Reg. No. 35,799

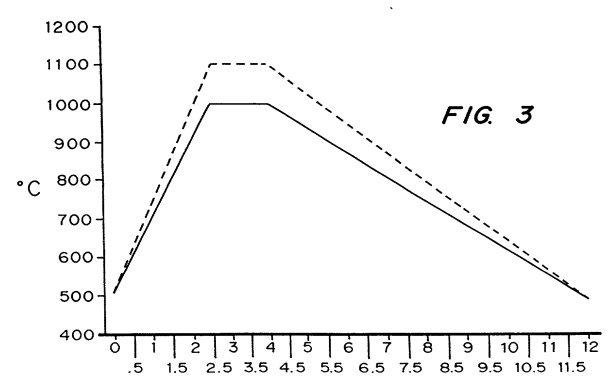
Attorney for Assignee

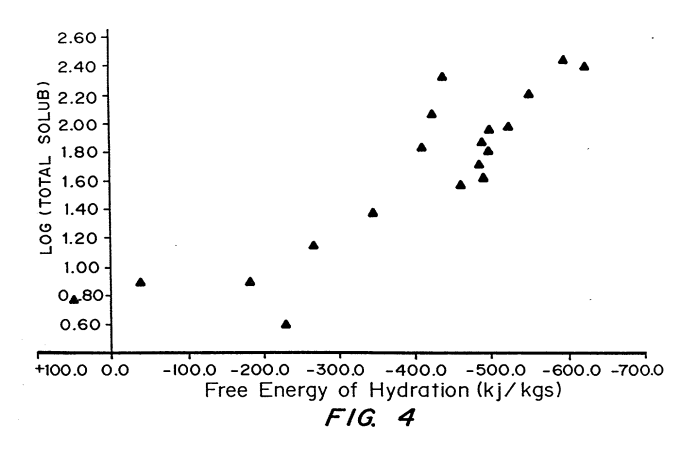
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404-815-6218
Attorney Docket No.: M8540/248465

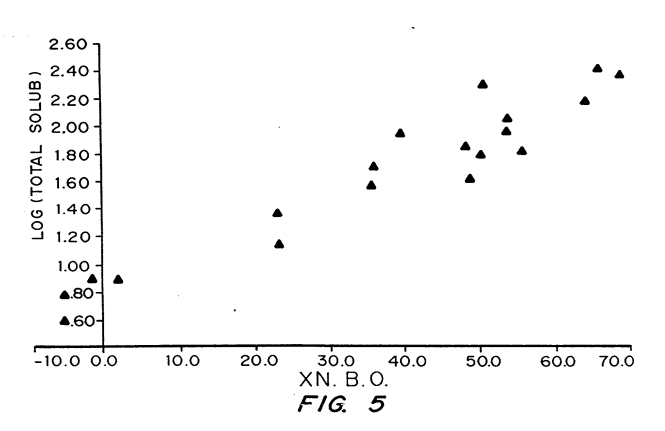
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COMBINED DECLARATION (Included Reference to PCT Inter	IBENED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY MB540/18831			
As a h	low named inventor, I bereby de	dare that:		
My residence, p	nt uffice address and citizenship	are as stated below ment to my a	Ame.	
Locat Strategic fr	e original, first and sole inventor. [Plaral names are listed below] of invention entitled:	(if only one name is listed below the subject matter which is cla) or an original, first and imed and for which a patent	
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	is attached hereia			
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	and was amended		,	
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	was filed as PCT international:	application Number		
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PRIOR FOREIGN/PCT	application(s) and any pr	JORITY CLAIMS UNDER 35	U.S.C. 119:	
COUNTRY (STEEL)	Application number	DATE OF FILING (day, possib, year)	PRIORITY CLADGED ENORR SETTIC 119	
GB GB	9200993.5	17 January 1992 (17.01.92)	⊠ ANY 0 MO	
GD.	9224612.3	24 November 1992 (24.11.92)	⊠ YES □NO	
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(Combined Declaration For Patent Application and Power of Attorney -- PTO 1391 [13-11]-page 1 of 2)

903D93 COMBINED DECLARATED WIR PATENT APPLICATION AND POWER OF ATTORNEY COMBRESD DECLARATION FOR PATRICT APPLICATION AND FOWER OF ATTORNEY (CONTINUED) (Includes Reference to PCT International Applications)

Attended to the same M8540/188317

I hereby claim the benefit under Title 35, United States Code, \$120 of any United States application(s) or PCT international application(s) designating the United States of Asseries that is/are listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in that/those prior application(s) in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application:

PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS DESIGNATING THE U.S. FOR BENEFIT UNDER 35 U.S.C. 120:

	U.S. APPLICATIONS			STATUS (Check One)		
U.S. APPLICATION NUMBER		U.S. FULING DATE	PATENTED	PENDING	ABANDONE	
08/536,5	87	28 September 1995 (28.69.95)			 	
0 8/899, 0	08/899,005 23 July 1997 (23.07.97)			X		
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APPLICATION NO	PCT FILING DATE	U.S. SERIAL NUMBERS ASSICNED (If my)				
PCT/GR93/ 00085	15 January 1993 (15.01.93)	08/039,486			×	
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POWER OF ATTORNEY: As a sensed inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (List name and registration number) Charles Y. Lackey 22,707; John M. Harrington 25,592; John S. Prutt 29,476; A. Jose Cartina 29,733; James L. Ewing. IV 30,630; Charles W. Callons 31,814; George T. Marcon 33,814; Dean W. Russell 31,452; Richard T. Peterson 35,320; Charles T. Simmons 35,359; Nora M. Tocups 35,717; Bruce D. Gray 35,799; Theodore R. Harper 35,896; Kristin L. Johnson 44,807; Geoff L. Sutchiffe 36,348; Pat Winston Kennedy 36,970; Mitchell G. Stuckwell 39,389; Michael J. Turton 40,852; Youths L. Kundupoglu 41,130; Benjamin D. Driscoll 41,571; Wilhurn L. Chesser 41,663; Alama G. Krisgman 41,747; J. Steven Gardner 41,772; Junes J. Bindseil 42,326; Richard H. Lilley 42,803; Edwins Thousas Washington 43,187; Camilla Camp Williams 43,992; Carl B. Massey P44,224; R. Whitney Winston P44,432; John William Ball, Jr. P44,433

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I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true, and further that these statements were made with the knowledge that willful fulse statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statement may jeopardize the validity of the application or any patent issuing thereon.

DONA Gary Anthony July

Jean-Louis Martin

903093 COMBINED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTOROGY

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

APPLICANTS:

Gary Anthony Jubb and Jean-

Louis Martin

Continuation of

GROUP ART

SERIAL NO.:

08/899,005

UNIT:

1755

DEPOSIT DATE:

March 4, 1999

EXAMINER:

Karl Group

FOR:

SALINE SOLUBLE INORGANIC FIBERS

Assistant Commissioner for Patents BOX PATENT APPLICATION

Attorney Docket No. 73405DIV

Washington, D.C. 20231

DATE: March 4, 1999

APPOINTMENT OF ASSOCIATE ATTORNEY

Sir:

Please recognize Bruce D. Gray, Reg. No. 35,799 of the firm of Kilpatrick Stockton LLP, 1100 Peachtree Street, Suite 2800, Atlanta, GA 30309-4530 as an Associate Attorney in the above-entitled application.

Respectfully submitted,

Dean W. Russell

Reg. No. 33,452

KILPATRICK STOCKTON LLP 1100 Peachtree Street, Suite 2800 Atlanta, GA 30309-4530 404.815.6500

ATLLIB01 741299 1

forming the basis for this application, divisional, or col	an 12 months from the filing date of this application is a PCT filing lication entering the United States as (1) the national stage, or (2) a nitinuation-in-part, then also complete ADDED PAGES TO COMBINED TTORNEY FOR DIVISIONAL, CONTINUATION OR CIP APPLICATION for benefit on(s) under 35 U.S.C. § 120
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	attorney(s) and/or agent(s) to prosecute this application Patent and Trademark Office connected therewith. (List
John S. Pratt, Reg. No. 2 James L. Ewing, IV, Reg. Patrea L. Pabst, Reg. No. Jamie Greene, Reg. No. 32 Cheryl K. Zalesky, Check th	No. 30,630 Charles T. Simmons, Reg. No. 31,284 Daniel M. Epstein, Reg. No. Claudia R. Adkison, Reg. No.
	s declaration and power of attorney is the authorization attorney(s) to accept and follow instructions from my
END CORRESPONDENCE TO	DIRECT TELEPHONE CALLS TO: (Name and telephone number)
John S. Pratt Kilpatrick & Cody 1100 Peachtree Street Suite 2800 Atlanta, Georgia 30309	Dean W. Russell (404) 815-6528
U.S.A.	DECLARATION
	ents made herein of my own knowledge are true and rmation and belief are believed to be true; and further
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(Rei 54-11/92 Pub 605)	FORM 1-1	1-11
(Declaration and Power	of Attorney [1-1]—page 4 o	f 5)
Full name of second joint	t inventor, if any <u>Jean-Louis N</u>	Martin
Jean-Louis	NMI	Martin
(GIVEN NAME) Inventor's signature	(MIDDLE INITIAL OR NAME)	FAMILY (OR LAST NAME)
Date <u>25/3/13</u>	Country of Citizenship _	France
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